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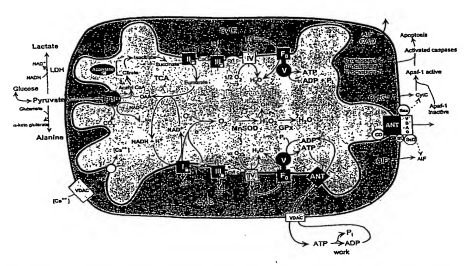
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[Continued on next page]

(54) Title: MITOCHONDRIAL BIOLOGY EXPRESSION ARRAYS



(57) Abstract: This invention provides a library of genes involved in mitochondrial biology, arrays containing probes for genes involved in mitochondrial biology, methods for making such arrays, and methods of using such arrays. Genes and probe sequences involved in mitochondrial biology in humans and mice are provided. The arrays of this invention are useful for determining mitochondrial biology gene expression profiles. Mitochondrial biology gene expression profiles are useful for determining expression profiles diagnostic of physiological conditions; diagnosing physiological conditions; identifying biochemical pathways, genes, and mutations involved in physiological conditions; identifying therapeutic agents useful for preventing and/or treating such physiological conditions; evaluating and/or monitoring the efficacy of such therapies, and creating and identifying animal models of human physiologic conditions. Arrays containing probes for all genes known to be involved in mitochondrial biology are provided, as well as arrays containing subsets of such probes.



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MITOCHONDRIAL BIOLOGY EXPRESSION ARRAYS

CROSS-REFERENCE TO RELATED APPLICATIONS

This application claims priority to U.S. Patent Application Serial No. 60/316,323 filed August 30, 2001, and to Canadian Patent Application Serial No. 2,356,540 filed August 31, 2001, both of which is hereby incorporated in their entirety by reference to the extent not inconsistent with the disclosure herein.

BACKGROUND OF THE INVENTION

Mitochondrial disorders are a complex and polygenic group of conditions with the patient's symptoms varying due to differences in energetic threshold effect of various tissues and the stochastic nature of mtDNA segregation. Consequently, most mitochondrial disorders are best classified by their genetic cause rather than a biochemical or phenotypic profile (Shoffner, J. M., and Wallace, D. C., (1995) "Oxidative phosphorylation diseases," In The Metabolic and Molecular Basis of Inherited Disease, C. R. Scriver, A. L. Beaudet, W. S. Sly and D. Valle, eds. (New York: McGraw-Hill), pp.1535-1609; Wallace, D. C., (1999) "Mitochondrial diseases in man and mouse" Science 283:1482-1488). Many mitochondrial diseases result from mutations in nuclear genes and a subset of these are known to act by destabilizing the mitochondrial genome. (Graham, B. et al., "A mouse model for mitochondrial myopathy and cardiomyopathy resulting from a deficiency in the heart/skeletal muscle isoform of the adenine nucleotide translocator," [1997] Nature Genetics 16:226-234; Shoffner, J. M., and Wallace, D.C., "Oxidative phosphorylation diseases. Disorders of two genomes," [1990] Advances in Human Genetics 19:267-330; Zhu, Z. et al., "SURF1, encoding a factor involved in the biogenesis of cytochrome c oxidase, is mutated in Leigh's syndrome" [1998] Nature Genetics 20:337-43).

The analysis of mitochondrial function in cultured cells using somatic cell genetics has been instrumental in the characterization of human mitochondrial disorders. Ethidium bromide and R-6G treatment have been used to create $\rho 0$ and mitochondria-less cell lines to analyze the maternal inheritance and biochemical phenotypes of many human mtDNA mutations (Chomyn, A. et al., "In vitro genetic transfer of protein synthesis and respiration defects to mitochondrial DNA-less cells with myopathy-patient mitochondria," [1991] Molecular and Cellular Biology 11:2236-2244; Jun, A.S. et al., "Use of transmitochondrial cybrids to assign a complex I defect to the mitochondrial DNA-encoded NADH

dehydrogenase subunit 6 gene mutation at nucleotide pair 14459 that causes Leber hereditary optic neuropathy and dystonia," [1996] Molecular and Cellular Biology 16:771-777; King, M. P. et al., "Defects in mitochondrial protein synthesis and respiratory chain activity segregate with the tRNA Leu(UUR) mutation associated with mitochondrial myopathy, encephalopathy, lactic acidosis, and stroke-like episodes," [1992] Molecular and Cellular Biology 12:480-490; Trounce, I. et al., "Cytoplasmic transfer of the mtDNA nt 8993 TG [ATP6] point mutation associated with Leigh's syndrome into mtDNA-less cells demonstrates cosegregation with a decrease in state III respiration and ADP/O ratio," [1994] Proc. Natl. Acad. Sci. U.S.A. 91:8334-8338). The creation of cybrid cell lines with identical nuclear backgrounds but different mtDNA genotypes allows the comparison of one mtDNA mutant to another without the potential interference of nuclear genome polymorphisms. These cybrid lines have generally been analyzed using biochemical techniques such as assaying cellular respiration or respiratory complex specific activities by enzymology. Some gene expression studies have been performed, but they have generally been done on single or small groups of genes (Heddi, A. et al., "Mitochondrial DNA expression in mitochondrial myopathies and coordinated expression of nuclear genes involved in ATP," [1993] J. Biological Chemistry 268:12156-12163; Heddi, A.et al., "Coordinate induction of energy gene expression in tissues of mitochondrial disease patients" [1999] J Biol Chem 274:22968-76).

Gene expression has been extensively studied. Although the regulation of mRNA abundance by changes in transcription or RNA degradation is by no means the only mechanism that regulates protein levels in a cell, virtually all differences in cell type or state can be correlated to changes in the mRNA abundance of several genes (Alizadeh, A.A. et al., "Distinct types of diffuse large B-cell lymphoma identified by gene expression profiling," [2000] Nature 403:503-11; DeRisi, J.L. et al., "Exploring the metabolic and genetic control of gene expression on a genomic scale," [1997] Science 278:680-686; Schena, M. et al., "Quantitative monitoring of gene expression patterns with a complementary DNA microarray" [1995] Science 270:467-70; Schena, M. et al., "Parallel human genome analysis: microarray-based expression monitoring of 1000 genes" [1996] Proc Natl Acad Sci USA 93:10614-9; Wallace D.C., grant abstract #2R01N502328-18; Kerstann, K.W. [2000] American Society for Human Genetics Abstract #1484; Kokoszka, J.E. [2000] American Society of Human Genetics Abstract #1618; Levy, S.E. [2001] American Society of Human Genetics Abstract #1501; Levy, S.E. [2000] "Genetic Alteration of the Mouse Mitochondrial

Genome and Effects on Gene Expression," Ph.D. Thesis, Emory University; Coskun, P.E. [2000] American Society of Human Genetics Abstract #1616; Sligh, J.E. [2000] American Society for Human Genetics Abstract #53; Murdock, D.G. [2000] American Society for Human Genetics Abstract #55; Levy S.E. [2000] Keystone Symposia Abstract 119; Wallace, D.C., Ellison Medical Foundation, Senior Scholar Award in Aging).

DNA microarray analysis has been used to study diffuse large B-cell lymphoma (DLBCL) where microarrays were used to expand the diagnosis of DLBCL (Alizadeh, A.A. et al., "Distinct types of diffuse large B-cell lymphoma identified by gene expression profiling," [2000] Nature 403:503-11). While standard histological and morphological techniques had defined subsets of DLBCL, array analysis revealed two clinically distinct classes. These two newly discovered classes were indistinguishable by standard pathology, but expression analysis showed a differential expression of hundreds of genes. Correlation of these molecular differences with differences in the progression of the disease and clinical outcome has revealed that these two classes of DLBCL could be considered separate diseases (Alizadeh, A.A. et al., "Distinct types of diffuse large B-cell lymphoma identified by gene expression profiling," [2000] Nature 403:503-11).

Mitochondrial DNA sequences have been associated with pathologies as described in U.S. patent numbers 5,670,320, 5,296,349, 5,185,244, and 5,494,794. Publications on the subject of mitochondrial biology include: Scheffler I.E. (1999) Mitochondria, Wiley-Liss, NY; Lestienne, P., Ed. (1999) Mitochondrial Diseases: Models and Methods, Springer-Verlag, Berlin; Methods in Enzymology (2000) 322: Section V Mitochondria and Apoptosis, Academic Press, CA; Mitochondria and Cell Death (1999) Princeton University Press, NJ; Papa S, Ferruciio G, and Tager J Eds. (1999) Frontiers of Cellular Bioenergetics: Molecular Biology, Biochemistry, and Physiopathology, Kluwer Academic / Plenum Publishers, NY; Lemasters J and Nieminen A (2001) Mitochondria in Pathogenesis, Kluwer Academic / Plenum Publishers, NY; MITOMAP, http://www.gen.emory.edu/cgi-gin/MITOMAP; Wallace D.C. (2001) "A mitochondrial paradigm for degenerative diseases and aging," Novartis Foundation Symposium 235:247-266; Wallace DC "Mitochondrial DNA in Aging and Disease" (August 1997) Scientific American 277:40-47; Wallace D.C. et al., "Mitochondrial biology, degenerative diseases and aging," (1998) BioFactors 7:187-190; Heddi, A. et al., "Coordinate Induction of Energy Gene Expression in Tissues of Mitochondrial Disease Patients" (1999) JBC 274:22968-22976; Wallace, D.C.

"Mitochondrial Diseases in Man and Mouse," (1999) Science 283:1482-1488; Saraste, M. "Oxidative Phosphorylation at the fin de siecle" (1999) Science 283:1488-1493; Kokoszka et. al., "Increased mitochondrial oxidative stress in the Sod2 (+/-) mouse results in the agerelated decline of mitochondrial function culminating in increased apoptosis," (2001) PNAS 98:2278-2283; Wallace, D.C. (2001) Mental Retardation and Developmental Disabilities 7:158-166; Wallace D.C. (2001) Am. J. Med. Gen. 106:71-93; and Wallace, D.C. (2001) EuroMit 5 Abstract.

The analysis of mitochondrial disorders has traditionally consisted of molecular and biochemical descriptions of the defect (Shoffner, J. M., and Wallace, D. C., (1995) "Oxidative phosphorylation diseases," In The Metabolic and Molecular Basis of Inherited Disease, C. R. Scriver, A. L. Beaudet, W. S. Sly and D. Valle, eds. (New York: McGraw-Hill), pp.1535-1609). Only a limited number of analyses of changes in oxidative phosphorylation (OXPHOS) genes expression have been performed in humans harboring mtDNA mutations (Heddi, A. et al., "Coordinate Induction of Energy Gene Expression in Tissues of Mitochondrial Disease Patients" (1999) *JBC* 274:22968-22976). The advent of mouse models for mitochondrial disease created by the inactivation of nuclear-encoded OXPHOS subunits has provided experimental material to study tissue-specific expression changes. (Murdock, D.G. et al., "Up-regulation of nuclear and mitochondrial genes in the skeletal muscle of mice lacking the heart/muscle isoform of the adenine nucleotide translocator," [1999] *J. Biol. Chem.* 274:14429-33.)

Nucleic acid arrays have been described, e.g., in patent nos. U.S. 5,837,832, U.S. 5,807,522, U.S. 6,007,987, U.S. 6,110,426, WO 99/05324, 99/05591, WO 00/58516, WO 95/11995, WO 95/35505A1, WO 99/42813, JP10503841T2, GR3030430T3, ES2134481T3, EP804731B1, DE69509925C0, CA2192095AA, AU2862995A1, AU709276B2, AT180570, EP 1066506, and AU 2780499. Such arrays can be incorporated into computerized methods for analyzing hybridization results when the arrays are contacted with prepared sample nucleotides, e.g., as described in PCT Publication WO 99/05574, and U.S. Patents 5,754,524; 6228,575; 5,593,839; and 5,856,101. Methods for screening for disease markers are also known to the art, e.g., as described in U.S. Patents 6,228,586; 6,160,104; 6,083,698; 6,268,398; 6,228,578; and 6,265,174.

All references cited herein are incorporated by reference in their entirety to the extent that they are not inconsistent with the disclosure herein. Citation of the above documents is not an admission that any of them are pertinent prior art.

SUMMARY OF THE INVENTION

This invention provides a library of genes involved in mitochondrial biology, arrays containing probes for genes involved in mitochondrial biology, methods for making such arrays, and methods of using such arrays. Genes and probe sequences involved in mitochondrial biology in humans and mice are provided. The arrays of this invention are useful for determining mitochondrial biology gene expression profiles. Mitochondrial biology gene expression profiles diagnostic of energy metabolism-related physiological conditions; diagnosing such physiological conditions; identifying biochemical pathways, genes, and mutations involved in such physiological conditions; identifying therapeutic agents useful for preventing and/or treating such physiological conditions; evaluating and/or monitoring the efficacy of such therapies; and creating and identifying animal models of human energy metabolism-related physiological conditions. Arrays containing probes for all genes known to be involved in mitochondrial biology are provided, as well as arrays containing subsets of such probes. The mitochondrial biology expression arrays of this invention contain probes of genes not previously recognized to participate in mitochondrial biology.

BRIEF DESCRIPTION OF THE DRAWINGS

FIG. 1 is a diagram of the mammalian mitochondrion showing mitochonrial energetics, and the relationship between energy production, reactive oxygen species (ROS) generation, and regulation of apoptosis.

FIG. 2 is a depiction of a hybridized mouse array of this invention. The picture of the hybridized array shows the image generated when the two channels representing the control or reference and experimental targets are overlaid. When viewed in color, the spots appear various shades of red, green and yellow. Red spots indicate a predominance of hybridization to control cDNAs, while green spots indicate the predominance of hybridization to the experimental target sample. Yellow spots indicate an equal hybridization of both samples. Spots that are yellow-green or orange when the array is shown in color are depicted as half yellow and green, or half red and yellow, respectively.

FIG. 3 shows the $\rho 0$ LMEB4 cell line gene expression scatter plot. The scatter plot shows the distribution of gene expression ratio for the $\rho 0$ LMEB4 sample. The diagonal dotted line indicates a ratio of 1 between the two samples. Any spot above the dotted line is up-regulated or more abundant in the $\rho 0$ LMEB4 experimental sample compared to the LM(TK) - control. Any spot below the dotted line is down-regulated or less abundant in the experimental sample compared to the control.

FIG. 4 shows NZB heart gene expression scatter plot. The scatter plot shows the distribution of gene expression ratio for the NZB heart tissue sample. The diagonal dotted line indicates a ratio of 1 between the two samples. Any spot above the dotted line is upregulated or more abundant in the NZB-mtDNA heart experimental sample compared to the "common" mtDNA control heart. Any spot below the dotted line is down-regulated or less abundant in the experimental sample compared to the control.

DETAILED DESCRIPTION OF THE INVENTION

An approach to examining the complex interaction between nuclear and cytoplasmic mitochondrial genes is through the use of arrays such as DNA arrays. DNA microarrays provide a means to profile the expression patterns of up to thousands of genes simultaneously, and knowing where and when a gene is expressed often provides insight into its biological function. The pattern of gene expression in a particular tissue or cell type can also provide detailed information about its state or condition.

Currently, DNA microarrays are the most efficient method to monitor correlative changes in gene expression and to investigate complex traits on a molecular level. Expression profiles assembled from multiple interrelated experiments are used to determine hierarchical connections between gene expression patterns underlying complex biological traits. These patterns are used to further define the molecular basis of complex disorders.

The mitochondrion is assembled from approximately 1000 protein-coding nuclear DNA (nDNA) and mitochondrial DNA (mtDNA) genes. Thirteen protein-coding mitochondrial genes are known, as shown in Table 1. The codon usage table of the mtDNA is known. It differs slightly from the universal code. For example, UGA codes for

tryptophan instead of termination, AUA codes for methionine instead of isoleucine, and AGA and AGG are terminators instead of coding for arginine.

Table 1

Gene	Map Locus ^a	Abbreviation	Location ^b
NADH dehydrogenase 1	MTND1	ND1	3307-4262
NADH dehydrogenase 2	MTND2	ND2	4470-5511
NADH dehydrogenase 3	MTND3	ND3	10059-10404
NADH dehydrogenase 4L	MTND4L	ND4L	10470-10766
NADH dehydrogenase 4	MTND4	ND4	10760-12137
NADH dehydrogenase 5	MTND5	ND5	12337-14148
NADH dehydrogenase 6	MTND6	ND6	14149-14673
Cytochrome b	MTCYB	Cytb	14747-15887
Cytochrome c oxidase I	MTCO1	COI	5904-7445
Cytochrome c oxidase II	MTCO2	COII	7586-8269
Cytochrome c oxidase III	MTCO3	COII	9207-9990
ATP synthase 6	MTATP6	ATP6	8527-9207
ATP synthase 8	MTATP8	ATP8	8366-8572

a,b As defined on MitoMap, http://www.gen.emory.edu/cgi-bin/MITOMAP, which is numbered relative to the Cambridge Sequence (Genbank accession no. J01415 and Andrews et al. (1999), A Reanalysis and Revision of the Cambridge Reference Sequence for Human Mitochondrial DNA, *Nature Genetics* 23:147.

As used herein "gene" refers to a unigene cluster, an expressed sequence, or a sequence that is transcribed and translated into a protein. Another word used in the art for "gene" is "locus." The National Institutes of Health (NIH) have instituted the term "unigene cluster" to refer to non-redundant sets of gene clusters. A stretch of DNA may be transcribed into several splice variants that share sequences, and these would be designated as belonging to one unigene cluster. As used herein "splice variant" refers to one version of several transcripts that are transcribed from one gene. As used herein "housekeeping gene" refers to a gene that is expressed at a similar level in almost all cell types.

As used herein "genes involved in mitochondrial biology" refers to mitochondrial genes and nuclear genes involved in cellular structures and functions such as intermediary metabolism, OXPHOS, mitochondrial transport, cellular bioeenergetics, cellular biogenesis,

cell cycle control, DNA replication, energy, metabolism, heat shock, stress, cellular matrix, cellular structural proteins, protein synthesis and translational control, signal transduction, transcription and transcriptional regulation, chromatin structure, reactive oxygen species (ROS) biology, and apoptosis.

"mtDNA" means mitochondrial DNA. "nDNA" means nuclear DNA.

As used herein "mitochondrial biology expression profile" refers to the expression patterns of genes involved in mitochondrial biology, such as is detected by probes derived from those genes, in a sample. The profile can be said to be of the sample or of the source from which the sample is derived. A profile may be measured independently, but a profile may also measured relative to a standard or control or other sample. A complete mitochondrial biology expression profile includes data on all genes known to be involved in mitochondrial biology for the species from which the sample is derived. The mitochondrial biology expression profile for a selected physiological condition is at least the expression pattern of genes determined to have altered expression diagnostic of that physiological condition, but the expression pattern of additional genes involved in mitochondrial biology may also be included.

As used herein "array" refers to an ordered set of isolated nucleic acid molecules or spots consisting of pluralities of substantially identical isolated nucleic acid molecules. Preferably the molecules are attached to a substrate. The spots or molecules are ordered so that the location of each (on the substrate) is known and the identity of each is known. Arrays on a micro scale can be called microarrays. Microarrays on solid substrates, such as glass or other ceramic slides, can be called gene chips or chips.

As used herein, an "isolated nucleic acid" is a nucleic acid outside of the context in which it is found in nature. An isolated nucleic acid is a nucleic acid the structure of which is not identical to that of any naturally occurring nucleic acid molecule. The term covers, for example: (a) a DNA which has the sequence of part of a naturally-occurring genomic DNA molecule but is not flanked by both of the coding or noncoding sequences that flank that part of the molecule in the genome of the organism in which it naturally occurs; (b) a nucleic acid incorporated into a vector or into the genomic DNA of a prokaryote or eukaryote in a manner such that the resulting molecule is not identical to any naturally-occurring vector or genomic

DNA; (c) a separate molecule such as a cDNA, a genomic fragment, a fragment produced by polymerase chain reaction (PCR), or a restriction fragment; and (d) a recombinant nucleotide sequence that is part of a hybrid gene, i.e., a gene encoding a fusion protein, or a modified gene having a sequence not found in nature.

As used herein "probe" refers to an isolated nucleic acid that is suitable for hybridizing to other nucleic acids when placed on a solid substrate. Probes for arrays can be as short as 20-30 nucleotides and up to as long as several thousand nucleotides. Probes can be single-stranded or double stranded. A probe usually comprises at least a partially known sequence that is used to investigate or interrogate the presence, absence, and/or amount of a complementing sequence. On the arrays of this invention, a probe is of such a sequence and the hybridization conditions of such stringency that each probe hybridizes substantially to only one type of nucleic acid per target sample.

As used herein, "target" or "target sample" refers to the collection of nucleic acids, e.g., reverse transcribed and labeled cDNA used as a prepared sample for array analysis. The target is interrogated by the probes of the array. A "target" or "target sample" may be a mixture of several prepared samples that are combined. For example, an experimental target sample may be combined with a differently labeled control sample and hybridized to an array, the combined samples being referred to as the "target" interrogated by the probes of the array. As used herein, "interrogated" means tested. Probes, targets, and hybridization conditions are chosen such that the probes are capable of interrogating the target, i.e., of hybridizing to complementary sequences in the target sample.

As used herein "physiological condition" refers to a healthy or unhealthy physiological state. As used herein "optimize an array for diagnosis" refers to selecting probes for an array such that only probes from genes necessary for diagnosis of one or more physiological conditions are included.

As used herein "printing" refers to the process of applying probes to a solid substrate, e.g., or applying arrays of probes to a solid substrate to make a gene chip. As used herein "glass slide" refers to a small piece of glass of the same dimensions as a standard microscope slide. As used herein, "prepared substrate" refers to a substrate that is prepared with a

substance capable of serving as an attachment medium for attaching the probes to the substrate, such as poly Lysine.

As used herein "selective hybridization" refers to hybridization at moderate to high stringency such that only sequences of an appropriate homology can remain bound. Selective hybridization is hybridization performed at stringency conditions such that probes only hybridize to target sample nucleic acids that they are intended to hybridize with. Depending on the sequences of the probes and the target, the hybridization conditions are chosen to be appropriately selective. For example, if human sequences are used as probes for interrogating a human sample, selective hybridization could be at high stringency because, allowing for neutral polymorphism in humans, the sequences would be about 99-100% identical. When applying a chimpanzee target prepared sample to an array containing human sequence probes, selective hybridization would be at a lower stringency. Since hybridizing a target to an array is performed at one chosen hybridization stringency, probes are chosen so that they can undergo selective hybridization with the appropriate target molecules at the same hybridization stringency. As used herein "homology" refers to nucleotide sequence identity to a sequence, a molecule, or its complement.

As used herein "mouse sample" refers to a sample derived from a mouse or a cell line derived from a mouse. Similarly, as used herein, "human sample" refers to a sample derived from a human or a cell line derived from a human. Samples preferably contain total RNA or messenger RNA (mRNA). As used herein "total RNA" refers to a combination of several types of RNA, including mRNA, from a cell or a group of cell. As used herein, "mRNA" refers to messenger RNA or RNA that has a 3' poly A tail. As used herein, a "prepared sample" or a "target" refers to a sample that has been labeled in preparation for array hybridization. A "prepared sample" or "target" is reverse transcribed and fluorescently labeled. As used herein "standard" refers to a sample or a dataset that is commonly used for comparison to unknown samples so that the unknown samples or datasets can be standardized for comparison to each other. As used herein, "control sample" and "reference sample" refer to samples that are used for comparison against an experimental sample.

As used herein, "clone" refers to an isolated nucleic acid molecule that may be stored in an organism such as *E. coli*. A clone is usually made of a vector and an insert. The insert usually contains a sequence of interest.

For mitochondrial diseases, the accuracy of current biochemical and phenotypic techniques has proven quite limited in distinguishing and diagnosing the various disorders. Recent technical and analytical advancements make it practical to analyze and quantitate the expression patterns of thousands of genes at once using arrays such as DNA microarrays. This invention applies these array techniques to the study of mitochondrial gene expression, in the design of specialized microarrays containing genes involved in mitochondrial biology. The arrays of this invention contain probes for genes not previously recognized to participate in mitochondrial biology.

Genes, or expressed sequences, involved in mitochondrial biology are involved in cellular structures and functions such as intermediary metabolism, OXPHOS, transport, cellular bioenergetics, cellular biogenesis, cell cycle control, DNA replication, energy, metabolism, heat shock, stress, cellular matrix, cellular structural proteins, protein synthesis and translational control, signal transduction, transcription and transcriptional regulation, chromatin structure, reactive oxygen species (ROS) biology and apoptosis. Alterations in mitochondrial functions are associated with a variety of physiological conditions including degenerative diseases. These functions are involved in many degenerative diseases. This invention provides a compilation of sequences involved in human and mouse mitochondrial biology.

The genes in the arrays of this invention were identified by a variety of techniques including searching databanks for sequences related to genes involved in processes similar to mitochondrial biology such as homologues of prokaryotic genes, and screening mitochondrial mutant cell lines and animal lines for genes having altered expression patterns. When a relevant gene was identified for one species, such as the mouse, the homologue for a second species, such as human, if known, was then included on the list of genes involved in mitochondrial biology for the second species. Mitochondrial mutant cell lines are cell lines that have at least one mutation in a gene involved in mitochondrial biology.

The microarrays or gene chips of this invention comprise probes placed in known positions on a solid substrate. A useful solid substrate is a specialized glass microscope slide. The arrays of this invention include arrays containing probes that detect some or all expressed sequences involved in mitochondrial biology in a selected species.

Arrays of this invention may contain control probes as well as probes for genes involved in mitochondrial biology. Controls that can be included on the arrays of this invention include hybridization controls and scanning controls. The controls can be positive or negative controls. One type of hybridization control is spotting the same probe for a gene involved in mitochondrial biology several times on one chip, each spot having different amounts of probe. This allows for the amount of probe of a given sequence to be optimized. Spotting too little probe may lead to a maximum hybridization signal resulting in a loss of data. Dimethyl sulfoxide (DMSO) can be used as a negative hybridization and scanning control. A spot of DMSO should give no signal. If there is any signal at a DMSO spot, the problem could be at hybridization or scanning steps. Plant sequences having sufficiently low homology with human and mouse sequences can be utilized as negative hybridization and scanning controls. Plant sequences should not give any signal. A signal at a plant spot could indicate a problem with hybridization, i.e. too low a hybridization stringency was used, or with scanning, i.e., the chip was inserted into the scanner at the incorrect orientation. Poly A can be used as a positive hybridization specificity/non specificity control. A poly A spot should always give intense hybridization. No signal at a poly A spot could be the result of use of too high a hybridization stringency. Cy3 or Cy5 incorporated into a PCR product can be a positive scanning control. A spot on an array of a PCR product, or any other nucleic acid, that includes fluorescent label, should always give a signal, and if this sequence has no homology with any other sequence in the target, there should only be a signal of the label included in the nucleic acid. Control probes and probes for genes involved in mitochondrial biology can be duplicated, triplicated, etc. on the chip as printing controls. Controls for arrays can be purchased from Stratagene (SpotReport TM , La Jolla, CA, USA).

Standard targets and reference targets are also useful with the arrays of this invention, as is known in the art. When a prepared sample target to be interrogated is applied to an array of this invention, the results of the test are measured, i.e. by scanning, and recorded. These results can be compared directly to other test results using a similar array. However, it is much more accurate to include a differently labeled standard target in the hybridization mix with the prepared sample target. The results of the experimental sample target are then standardized, so that they can be compared accurately to the results of hybridizations of other sample targets. If ten different prepared sample targets are hybridized to arrays of this invention, simultaneously with the same prepared standard target, then the results of the ten

sample targets can be accurately compared to each other. A prepared reference or control target for comparison can also be particularly pertinent to the experiment being performed. A prepared reference target could be a target sample derived from the same cell type from an animal of the same sex, age, and nuclear background as the experimental target sample, except for one difference, such as a different phenotype or treatment. Comparing the results of the experimental target with the results of an appropriate reference target yields a profile associated with the one difference being tested. When the hybridization results of a first sample are compared to the hybridization results of a second sample, the comparison can occur while the hybridization results of the first sample are being measured and recorded, or afterwards, by comparing the measured and recorded hybridization results of the two samples.

Probes on an array may be as short as about 20-30 nucleotides long or as long as the entire gene or clone from which they are derived, which may be up to several kilobases. A probe sequence may be identical (have 100% homology) to the portion of the gene it hybridizes to or it may be a mutated sequence. Mutated probes have less than 100% homology, such as about 98% homology, about 95% homology, about 90% homology, about 80% homology, or about 75% homology, or less, with the portions of the genes to which they hybridize. Arrays are designed such that all probes on an array can hybridize to their corresponding genes at about the same hybridization stringency. Probes for arrays used for interrogating samples usually do not contain sequences such as repetitive sequences that would hybridize substantially with nucleic acids derived from more than one gene, i.e., transcripts or cDNAs. Probes for arrays should be unique at the hybridization stringencies used. Statistically, to be unique in the total human genome, probes should be at least about fifteen nucleotides long. A unique probe is only able to hybridize with one type of nucleic acid per target. A probe is not unique if at the hybridization stringency used, it hybridizes with nucleic acids derived from two different genes, i.e. related genes. The homology of the sequence of the probe to the gene and the hybridization stringency used help determine whether a probe is unique when testing a selected sample. Probes also may not hybridize with different nucleic acids derived from the same gene, i.e., splice variants. The location in the gene of the sequence used for the probe also helps determines whether a probe is unique when testing a selected sample. If the splice variants of a gene are known, ideally several different probes sequences are chosen from that gene for an array, such that each probe can only hybridize to nucleic acid derived from one of the splice variants. References for

sequences of probes useful for arrays of this invention are compiled in Tables 3-5 and in the sequence listings. Other equivalent probes derived from the gene sequences from which the Tables 3-5 probes are derived, are also useful for the arrays of this invention. Arrays of this invention are used at hybridization conditions allowing for selective hybridization. At conditions of selective hybridization, probes hybridize with nucleic acid from only one gene. When an array is simultaneously hybridized with two targets or two prepared samples, each probe may hybridize with a nucleic acid in each prepared sample or target. When these two nucleic acids are from the same unigene cluster, the probe is said to hybridize with one gene, despite the fact that these nucleic acids may contain different labels.

Sequences of genes involved in mitochondrial biology from other species can be used to make probes that are useful in the arrays of this invention as long as they hybridize at about the same hybridization stringency as other probes on an array. Sequences that are only able to hybridize at a substantially lower stringency, such as plant sequences, are useful as negative controls.

The arrays of this invention can be utilized to determine profiles for related species by modifying the hybridization stringency appropriately. Sequence homology between organisms is known in the art. For example, human and chimpanzee sequences are about 98% identical. Consequently, human arrays are useful for profiling chimpanzees, with an appropriate lowering of the hybridization stringency. Hybridization stringency can be lowered by modifying hybridization components such as salt concentrations and hybridization and/or wash temperatures, as is known in the art.

The sequences useful for the arrays of this invention are useful for designing arrays for other species as well. To create an array for a new organism, the known sequences from the new organism, including expressed sequence tags (ESTs), are compared, by methods known to the art, with the sequences known to already be useful for other mitochondrial biology arrays. Sequence comparisons may be performed at the nucleic acid or polypeptide level. Homologous and analogous sequences from the new organism are thereby identified and selected for the new organism's mitochondrial array. The probes on the arrays of this invention are also useful as probes for identifying candidates for the new organism's array using molecular biology techniques that are standard in the art such as screening libraries.

All sequences given herein are meant to encompass the complementary strand, as well as double-stranded polynucleotides comprising the given sequence.

Microarrays of this invention can contain as few as two probes to as many as all the probes diagnostic of the selected physiological condition to be tested. Microarrays of this invention may also contain probes for all genes involved in mitochondrial biology. The arrays of this invention may contain probes for at least about five genes, at least about ten genes, at least about twenty-five genes, at least about fifty genes, at least about 100 genes, at least about 500 genes, or at least about 1000 genes. The mouse array may contain probes for at least about 950 genes and the human array may contain probes for at least about 600 genes. Arrays of this invention may comprise more than about five spots, more than about ten spots, more than about twenty-five spots, more than about one hundred spots, more than about 500 spots, or more than about 1000 spots.

Using microarrays may require amplification of target sequences (generation of multiple copies of the same sequence) of sequences of interest, such as by PCR or reverse transcription. As the nucleic acid is copied, it is tagged with a fluorescent label that emits light like a light bulb. The labeled nucleic acid is introduced to the microarray and allowed to react for a period of time. This nucleic acid sticks to, or hybridizes, with the probes on the array when the probe is sufficiently complementary to the labeled, amplified, sample nucleic acid. The extra nucleic acid is washed off of the array, leaving behind only the nucleic acid that has bound to the probes. By obtaining an image of the array with a fluorescent scanner and using software to analyze the hybridized array image, it can be determined if, and to what extent, genes are switched on and off, or whether or not sequences are present, by comparing fluorescent intensities at specific locations on the array. The intensity of the signal indicates to what extent a sequence is present. In expression arrays, high fluorescent signals indicate that many copies of a gene are present in a sample, and lower fluorescent signal shows a gene is less active. By selecting appropriate hybridization conditions and probes, this technique is useful for detecting single nucleotide polymorphisms (SNPs) and for sequencing. Methods of designing and using microarrays are continuously being improved (Relogio, A. et al. (2002) Nuc. Acids. Res. 30(11):e51; Iwasaki, H et al. (2002) DNA Res. 9(2):59-62; and Lindroos, K. et al. (2002) Nuc. Acids. Res. 30(14):E70).

Arrays of this invention may be made by any array synthesis methods known in the art such as spotting technology or solid phase synthesis. Preferably the arrays of this invention are synthesized by solid phase synthesis using a combination of photolithography and combinatorial chemistry. Some of the key elements of probe selection and array design are common to the production of all arrays. Strategies to optimize probe hybridization, for example, are invariably included in the process of probe selection. Hybridization under particular pH, salt, and temperature conditions can be optimized by taking into account melting temperatures and by using empirical rules that correlate with desired hybridization behaviors. Computer models may be used for predicting the intensity and concentration-dependence of probe hybridization.

Arrays, also called DNA microarrays or DNA chips, are fabricated by high-speed robotics, generally on glass but sometimes on nylon substrates, for which probes (Phimister, B. (1999) Nature Genetics 21s:1-60) with known identity are used to determine complementary binding. An experiment with a single DNA chip can provide researchers information on thousands of genes simultaneously. There are several steps in the design and implementation of a DNA array experiment. Many strategies have been investigated at each of these steps: 1) DNA types; 2) Chip fabrication; 3) Sample preparation; 4) Assay; 5) Readout; and 6) Software (informatics).

There are two major application forms for the array technology: 1) Determination of expression level (abundance) of genes; and 2) Identification of sequence (gene / gene mutation). There appear to be two variants of the array technology, in terms of intellectual property, of arrayed DNA sequence with known identity: Format I consists of probe cDNA (500~5,000 bases long) immobilized to a solid surface such as glass using robot spotting and exposed to a set of targets either separately or in a mixture. This method, "traditionally" called DNA microarray, is widely considered as having been developed at Stanford University. (R. Ekins and F.W. Chu "Microarrays: their origins and applications," [1999] Trends in Biotechnology, 17:217-218). Format II consists of an array of oligonucleotide (20~80-mer oligos) or peptide nucleic acid (PNA) probes synthesized either in situ (on-chip) or by conventional synthesis followed by on-chip immobilization. The array is exposed to labeled sample DNA, hybridized, and the identity/abundance of complementary sequences is determined. This method, "historically" called DNA chips, was developed at Affymetrix, Inc., which sells its photolithographically fabricated products under the GeneChip®

trademark. Many companies are manufacturing oligonucleotide-based chips using alternative *in-situ* synthesis or depositioning technologies.

Probes on arrays can be hybridized with fluorescently-labeled target polynucleotides and the hybridized array can be scanned by means of scanning fluorescence microscopy. The fluorescence patterns are then analyzed by an algorithm that determines the extent of mismatch content, identifies polymorphisms, and provides some general sequencing information (M. Chee et al., [1996] *Science* 274:610). Selectivity is afforded in this system by low stringency washes to rinse away non-selectively adsorbed materials. Subsequent analysis of relative binding signals from array elements determines where base-pair mismatches may exist. This method then relies on conventional chemical methods to maximize stringency, and automated pattern recognition processing is used to discriminate between fully complementary and partially complementary binding.

Devices such as standard nucleic acid microarrays or gene chips, require data processing algorithms and the use of sample redundancy (i.e., many of the same types of array elements for statistically significant data interpretation and avoidance of anomalies) to provide semi-quantitative analysis of polymorphisms or levels of mismatch between the target sequence and sequences immobilized on the device surface. Such algorithms and software useful for statistical analysis are known to the art.

Using microarrays first requires amplification (generation of multiple copies of the same gene) of genes of interest, such as by reverse transcription. As the nucleic acid is copied, it is tagged with a fluorescent label that emits light like a light bulb. The labeled nucleic acid is introduced to the microarray and allowed to react for a period of time. This nucleic acid sticks to, or hybridizes, with the probes on the array when the probe is sufficiently complementary to the nucleic acid in the prepared sample. The extra nucleic acid is washed off of the array, leaving behind only the nucleic acid that has bound to the probes. By obtaining an image of the array with a fluorescent scanner and using software to analyze the hybridized array image, it can be determined if and to what extent genes are switched on and off, or whether or not sequences are present, by comparing fluorescent intensities at specific locations on the array. High fluorescent signals indicate that many copies of a gene are present in a prepared sample, and lower fluorescent signal shows a gene is less active. Expression levels for various genes under different conditions can be directly compared, such

as for a cancer cell and a normal cell. Similarly, it can be determined what genes are turned on and off in response to certain stimuli such as a drug. Such information is valuable because it identifies genes in disease pathways and also is predictive of either efficacy or toxicity of drugs.

Detecting a particular polymorphism can be accomplished using two probes. One probe is designed to be perfectly complementary to a target sequence, and a partner probe is generated that is identical except for a single base mismatch in its center. In the Affymetrix system, these probe pairs are called the Perfect Match probe (PM) and the Mismatch probe (MM). They allow for the quantitation and subtraction of signals caused by non-specific cross-hybridization. The difference in hybridization signals between the partners, as well as their intensity ratios, serve as indicators of specific target abundance, and consequently of the sequence.

Arrays can rely on multiple probes to interrogate individual nucleotides in a sequence. The identity of a target base can be deduced using four identical probes that vary only in the target position, each containing one of the four possible bases. Alternatively, the presence of a consensus sequence can be tested using one or two probes representing specific alleles. To genotype heterozygous or genetically mixed samples, arrays with many probes can be created to provide redundant information, resulting in unequivocal genotyping.

Probes fixed on solid substrates and targets (nucleotide sequences in the sample) are combined in a hybridization buffer solution and held at an appropriate temperature until annealing occurs. Thereafter, the substrate is washed free of extraneous materials, leaving the nucleic acids on the target bound to the fixed probe molecules allowing for detection and quantitation by methods known in the art such as by autoradiograph, liquid scintillation counting, and/or fluorescence. As improvements are made in hybridization and detection techniques, they can be readily applied by one of ordinary skill in the art. As is well known in the art, if the probe molecules and target molecules hybridize by forming a strong noncovalent bond between the two molecules, it can be reasonably assumed that the probe and target nucleic acid are essentially identical, or almost completely complementary if the annealing and washing steps are carried out under conditions of high stringency. The detectable label provides a means for determining whether hybridization has occurred.

When using oligonucleotides or polynucleotides as hybridization probes, the probes may be labeled. In arrays of this invention, the target may instead be labeled by means known to the art. Target may be labeled with radioactive or non-radioactive labels. Targets preferably contain fluorescent labels.

Various degrees of stringency of hybridization can be employed. The more stringent the conditions are, the greater the complementarity that is required for duplex formation. Stringency can be controlled by temperature, probe concentration, probe length, ionic strength, time, and the like. Hybridization experiments are often conducted under moderate to high stringency conditions by techniques well know in the art, as described, for example in Keller, G.H., and M.M. Manak (1987) DNA Probes, Stockton Press, New York, NY., pp. 169-170, hereby incorporated by reference. However, sequencing arrays typically use lower hybridization stringencies, as is known in the art.

Moderate to high stringency conditions for hybridization are known to the art. An example of high stringency conditions for a blot are hybridizing at 68° C in 5X SSC/5X Denhardt's solution/0.1% SDS, and washing in 0.2X SSC/0.1% SDS at room temperature. An example of conditions of moderate stringency are hybridizing at 68° C in 5X SSC/5X Denhardt's solution/0.1% SDS and washing at 42° C in 3X SSC. The parameters of temperature and salt concentration can be varied to achieve the desired level of sequence identity between probe and target nucleic acid. See, e.g., Sambrook et al. (1989) vide infra or Ausubel et al. (1995) Current Protocols in Molecular Biology, John Wiley & Sons, NY, NY, for further guidance on hybridization conditions.

The melting temperature is described by the following formula (Beltz, G.A. et al., [1983] Methods of Enzymology, R. Wu, L. Grossman and K. Moldave [Eds.] Academic Press, New York 100:266-285).

Tm=81.50 C + 16.6 Log[Na+]+0.41(+G+C)-0.61(%formamide)-600/length of duplex in base pairs.

Washes can typically be carried out as follows: twice at room temperature for 15 minutes in 1X SSPE, 0.1% SDS (low stringency wash), and once at TM-200 C for 15 minutes in 0.2X SSPE, 0.1% SDS (moderate stringency wash).

Nucleic acid useful in this invention can be created by Polymerase Chain Reaction (PCR) amplification. PCR products can be confirmed by agarose gel electrophoresis. PCR is a repetitive, enzymatic, primed synthesis of a nucleic acid sequence. This procedure is well known and commonly used by those skilled in this art (see Mullis, U.S. Patent Nos. 4,683,195, 4,683,202, and 4,800,159; Saiki et al. [1985] Science 230:1350-1354). PCR is used to enzymatically amplify a DNA fragment of interest that is flanked by two oligonucleotide primers that hybridize to opposite strands of the target sequence. The primers are oriented with the 3' ends pointing towards each other. Repeated cycles of heat denaturation of the template, annealing of the primers to their complementary sequences, and extension of the annealed primers with a DNA polymerase result in the amplification of the segment defined by the 5' ends of the PCR primers. Since the extension product of each primer can serve as a template for the other primer, each cycle essentially doubles the amount of DNA template produced in the previous cycle. This results in the exponential accumulation of the specific target fragment, up to several million-fold in a few hours. By using a thermostable DNA polymerase such as the Taq polymerase, which is isolated from the thermophilic bacterium Thermus aquaticus, the amplification process can be completely automated. Other enzymes that can be used are known to those skilled in the art.

Polynucleotide sequences of the present invention can be truncated and/or mutated such that certain of the resulting fragments and/or mutants of the original full-length sequence can retain the desired characteristics of the full-length sequence. A wide variety of restriction enzymes that are suitable for generating fragments from larger nucleic acid molecules are well known. In addition, it is well known that Bal31 exonuclease can be conveniently used for time-controlled limited digestion of DNA. See, for example, Maniatis (1982) Molecular Cloning: A Laboratory Manual, Cold Spring Harbor Laboratory, New York, pages 135-139, incorporated herein by reference. See also Wei et al. (1983) J. Biol. Chem. 258:13006-13512. By use of Bal31 exonuclease (commonly referred to as "erase-a-base" procedures), the ordinarily skilled artisan can remove nucleotides from either or both ends of the subject nucleic acids to generate a wide spectrum of fragments that are functionally equivalent to the subject nucleotide sequences. One of ordinary skill in the art can, in this manner, generate hundreds of fragments of controlled, varying lengths from locations all along the original molecule. The ordinarily skilled artisan can routinely test or screen the generated fragments for their characteristics and determine the utility of the

fragments as taught herein. It is also well known that the mutant sequences can be easily produced with site-directed mutagenesis. See, for example, Larionov, O.A. and Nikiforov, V.G. (1982) *Genetika* 18(3):349-59; and Shortle, D. et al., (1981) *Annu. Rev. Genet.* 15:265-94, both incorporated herein by reference. The skilled artisan can routinely produce deletion, insertion, or substitution-type mutations and identify those resulting mutants that contain the desired characteristics of wild-type sequences, or fragments thereof.

Thus, mutational, insertional, and deletional variants of the disclosed nucleotide sequences can be readily prepared by methods which are well known to those skilled in the art. These variants can be used in the same manner as the exemplified primer sequences so long as the variants have substantial sequence homology with the original sequence. As used herein, substantial sequence homology refers to homology that is sufficient to enable the variant polynucleotide to function in the same capacity as the polynucleotide from which the probe was derived. Homology is greater than 80%, greater than 85%, greater than 90%, or greater than 95%. The degree of homology or identity needed for the variant to function in its intended capacity depends upon the intended use of the sequence. It is well within the skill of a person trained in this art to make mutational, insertional, and deletional mutations that are equivalent in function or are designed to improve the function of the sequence or otherwise provide a methodological advantage.

Percent sequence identity of two nucleic acids may be determined using the algorithm of Karlin and Altschul (1990) *Proc. Natl. Acad. Sci. USA* 87:2264-2268, modified as in Karlin and Altschul (1993) *Proc. Natl. Acad. Sci. USA* 90:5873-5877. Such an algorithm is incorporated into the NBLAST and XBLAST programs of Altschul et al. (1990) *J. Mol. Biol.* 215:402-410. BLAST nucleotide searches are performed with the NBLAST program, score = 100, wordlength = 12, to obtain nucleotide sequences with the desired percent sequence identity. To obtain gapped alignments for comparison purposes, Gapped BLAST is used as described in Altschul et al. (1997) *Nucl. Acids. Res.* 25:3389-3402. When utilizing BLAST and Gapped BLAST programs, the default parameters of the respective programs (NBLAST and XBLAST) are used. See http://www.ncbi.nih.gov.

Standard techniques for cloning, DNA isolation, amplification and purification, for enzymatic reactions involving DNA ligase, DNA polymerase, restriction endonucleases and the like, and various separation techniques useful herein are those known and commonly

employed by those skilled in the art. A number of standard techniques are described in Sambrook et al. (1989) Molecular Cloning, Second Edition, Cold Spring Harbor Laboratory, Plainview, New York; Maniatis et al. (1982) Molecular Cloning, Cold Spring Harbor Laboratory, Plainview, New York; Wu (ed.) (1993) Meth. Enzymol. 218, Part I; Wu (ed.) (1979) Meth. Enzymol. 68; Wu et al. (eds.) (1983) Meth. Enzymol. 100 and 101; Grossman and Moldave (eds.) Meth. Enzymol. 65; Miller (ed.) (1972) Experiments in Molecular Genetics, Cold Spring Harbor Laboratory, Cold Spring Harbor, New York; Old and Primrose (1981) Principles of Gene Manipulation, University of California Press, Berkeley; Schleif and Wensink (1982) Practical Methods in Molecular Biology; Glover (Ed.) (1985) DNA Cloning Vol. I and II, IRL Press, Oxford, UK; Hames and Higgins (Eds.) (1985) Nucleic Acid Hybridization, IRL Press, Oxford, UK; Setlow and Hollaender (1979) Genetic Engineering: Principles and Methods, Vols. 1-4, Plenum Press, New York; and Ausubel et al. (1992) Current Protocols in Molecular Biology, Greene/Wiley, New York, NY.

Abbreviations and nomenclature, where employed, are deemed standard in the field and commonly used in professional journals such as those cited herein.

The arrays of this invention are useful for defining expression signatures or profiles for mitochondrial diseases, as well as distinguishing clinical disorders that result from OXPHOS dysfunction, oxidative stress, apoptosis, and aging. The microarrays of this invention are useful for providing profiles for whole classes of mitochondrial diseases that have common underlying pathophysiological mechanisms. The data obtained from using these arrays are useful in the identification of pathways involved in these diseases and in the design of efficient therapies for treating these diseases.

The arrays of this invention are useful for determining mitochondrial biology expression profiles and for sample evaluation using those profiles. The arrays of this invention are useful for diagnosis, for identifying pathways, genes, and mutations involved in physiological conditions, for creating animal models of human physiological conditions, and for designing curative and preventative therapies and evaluating their effectiveness.

The arrays of this invention are useful for determining mitochondrial biology expression profiles of organisms, such as humans, mice, and closely related species; tissues and organs of such organisms; cell types of such organisms; and cell lines derived from such organisms. An individual can be tested at any age, including as a fetus, neonate, infant, child,

adolescent, mature adult, senior, and deceased. Using standard targets, the arrays of this invention are useful for comparing mitochondrial biology profiles of different individuals or cells.

The arrays of this invention are useful for determining the profile associated with a physiological condition such as an energy-metabolism-related physiological condition. Physiological conditions can be healthy conditions or pathological conditions. Examples of healthy conditions in humans are centenaria and physical fitness. An example of a pathological condition in humans is Leigh's syndrome (LS). By determining profiles from individuals, with and without such physiological conditions, and comparing them, the mitochondrial biology profile representative and descriptive of the physiological condition can be determined, such as for humans in Examples 4-5. Profiles can similarly be determined for cells lines with phenotypes or genotypes associated with physiological conditions, such as in Examples 13-15. Profiles can also be determined for non-human animals, including mouse strains, with physiological conditions as in Examples 8-12, 16, and 19. The arrays of this invention are useful for determining the range of normal variation of expression of genes involved in mitochondrial biology, as in Example 20. When the arrays of this invention are used to determine a profile associated with a physiological condition, prepared target samples or pooled prepared target samples, of individuals with and without the physiological condition, but otherwise similar, are hybridized to an array of this invention. The hybridization of the prepared samples are measured and compared to, if possible, determine a profile associated with the physiological condition. The profile may be optimized by statistical analysis, as is known in the art, to only contain profile data on probes necessary for diagnosing the physiological condition.

The profile associated with a physiological condition can then be used for diagnosis or evaluation using the arrays of this invention, such as in Example 7. The profile of the physiological condition can be analyzed and the analysis used to optimize an array for diagnosis of the physiological condition. An optimized array for diagnosis of a physiological condition minimally contains at least one probe for the one or more genes that have altered expression levels in the context of the physiological condition, and probes for enough genes to eliminate other likely diagnoses. Diagnosis involves collecting a sample from an individual who might have the physiological condition, and determining the profile of the prepared sample using an array of this invention, using an array containing probes for all

genes involved in mitochondrial biology or fewer probes with at least as many probes as necessary for an array optimized for diagnosis of the physiological condition. The profile of the individual is then compared to the profile of the physiological condition, and the comparison is analyzed to determine the likelihood that the individual has the physiological condition. Arrays of this invention can also be used for screening individuals who are not suspected of having the particular physiological condition. A sample is collected from such an individual, prepared, and the mitochondrial biology profile of the individual is determined using an array of this invention, e.g., an array containing probes for all genes involved in mitochondrial biology. The profile of this individual is then compared to known mitochondrial biology profiles of one or more physiological conditions that the individual may have, to determine if the profile of the individual is indicative of a diagnosable physiological condition. As demonstrated in Example 16, the arrays of this invention are also useful for detecting profiles indicative of physiological conditions before the appearance of other symptoms.

The profile of, or associated with, a physiological condition is also useful for identifying biochemical pathways affected by the physiological condition and genes involved in causation of the physiological condition. If a profile of a physiological condition demonstrates alteration in the expression of a gene, that gene is a candidate for sequencing to identify a mutation causing the physiological condition. If a profile demonstrates alteration of expression of several genes, then genes known to regulate those are candidates for sequencing to identify a mutation causing the physiological condition. Example 3 describes using the arrays of this invention for the identification of mutations associated with physiological conditions.

The profile of a physiological condition is useful for creating and/or identifying animal models of human physiological conditions. The profile of a physiological condition may suggest types of mutations, such as knockouts, to create in order to mimic the physiological condition in an animal. The arrays of this invention are also useful for screening genetically engineered or other mutated populations to identify an individual animal having a similar profile, and thus associated with the physiological condition.

The same individual can be profiled, using arrays of this invention, repeatedly over time or after exposure to various environmental conditions, thereby determining the effects of

time or exposure. Equivalent individuals can also be profiled, using the arrays of this invention, at different ages or after exposure to different environmental conditions, thereby determining the effects of time or exposure. For example, a control group of mice of a particular genotype and of a particular age can be compared, using the arrays of this invention, to a group of experimental mice of the same genotype and age, that has been exposed to a certain environmental hazard, to determine the effects of the environmental hazard. Cell lines, as well as organisms, can be profiled after exposure to different environmental conditions, as in Example 15. Arrays of this invention are also useful for determining the effects of aging. Examples 8 and 19 demonstrate differences in profiles at different ages.

Therapy is an environmental condition, the effects of which can be tested using the arrays of this invention. Identification of the pathways affected in a physiological condition allows identification of therapies useful to treat individuals having the physiological condition. For example, if profiles are determined for the effects of classes of therapeutic agents, as new physiological conditions are profiled, relevant therapeutic agents can be easily identified. The profile of a physiological condition is useful for testing candidate therapies for treating individuals with the physiological condition. An individual, with or without the physiological condition, an animal model of the physiological condition in humans, or a cell line representative of an individual with the physiological condition, can be treated with a candidate therapy. A sample for profiling is collected after treatment, prepared, the profile is determined using an array of this invention, and compared to the profile of the same individual before treatment or to equivalent individuals or cells without treatment to determine the effect of the treatment. Therapies reversing the effects of the physiological condition can thereby be identified. Preventative therapies and therapies causing desired physiological conditions can similarly be identified.

The arrays of this invention are useful for monitoring the effectiveness of a therapy for a particular individual as well as for a population. The profile of a diagnosed individual can be determined, the individual given a therapy, and then the profile of the individual determined again, using the arrays of this invention. The therapy can be modified and the profile retested, until a satisfactory treated profile is obtained.

Arrays containing probes hybridizing at moderate to high stringency with human genes involved in mitochondrial biology are used for assaying prepared samples from humans, human cell lines, and prepared samples from closely related species. Arrays containing probes hybridizing at moderate to high stringency with mouse genes involved in mitochondrial biology are used for assaying prepared samples from mice, mouse cell lines, and prepared samples from closely related species.

The arrays of this invention are made using probes for genes involved in mitochondrial biology. Probes can be selected and generated from the lists of clones and sequences in Tables 3-5, or from sequences and clones representing genes involved in mitochondrial biology not listed in these tables. Probes can be generated in vitro by nucleic acid synthesis, PCR, cloning techniques or other techniques known in the art. Flanking or vector sequence may be minimized in the probe. Probes generated from Research Genetics clones (ResGen/Invitrogen, Carlsbad, CA) can be amplified by PCR as described in Example 22. Optionally, control probes are also selected for the arrays of this invention. Examples of clones and sequences for making control probes are listed in Table 6, SEQ ID NOS:3041-3044. If housekeeping genes are chosen as positive controls, usually they are derived from the same species as the non-control probes. Housekeeping gene probes are available from Stratagene (Spot ReportTM, La Jolla, CA, USA).

Examples of housekeeping genes are shown in Table 2. Housekeeping genes generally have a consistent amount of expression in all cells. Using the arrays of this invention, the expression of the 25 housekeeping genes listed in Table 2 were compared in 4 cell lines, LMEB4, NZB, 501-1, and the LM(TK) - cell line grown in media supplemented with glucose, pyruvate, and uridine (GUP). Some variability was present between cell lines. Housekeeping genes were also tested in 6 different mouse tissue samples (brain, heart, liver, kidney, spleen and muscle) in two strains of mice, CAP^R and NZB. Variation was again present, but slight.

Table 2

Description	Functional Class
Actin-gamma	Structural gene
A272 Capping protein	Structural gene
Glyceraldehyde phosphate dehydrogenase	Metabolism-glycolysis
DNA ligase I	DNA repair/synthesis
β-actin	Structural
Alkaline Phosphatase	Unclassified
40s Ribosomal protein S15	Protein synthesis
Hypozanthine phosphoribosyl transferase (HPRT)	Metabolism-nucleotide
Ribosomal protein L15	Protein synthesis
Ribosomal Protein S29	Protein translation
Acient ubiquinating protein	Metabolism-protein
Glyceraldehyde 3-phosphate dehydrogenase	Metabolism-glycolysis
Actin-α (skeletal muscle)	Structural
Murine ornithine decarboxylase	Metabolism-amino acid
calcium binding protein Cab45	Calcium homeostasis
Ribosomal protein L1A	Protein synthesis
RNA splicing protein	RNA processing
Actin-gamma (smooth muscle)	Structural gene
E2F transcription factor	Transcriptional regulation
Ubiquitin	Unclassified
Myosin 1	Structural
HPRT	Metabolism-nucleotide
Phospholipase A2 (14-3-3 zeta/delta)	Signal transduction
HPRT	Metabolism-nucleotide
Ribosomal protein L3	Protein synthesis

Arrays can be printed on solid substrates, e.g., glass microscope slides. Before printing, slides are prepared to provide a substrate for binding as in Example 23. Arrays can be printed using any printing techniques and machines known in the art. Printing involves placing the probes on the substrate, attaching the probes to the substrate, and blocking the substrate to prevent non-specific hybridization, as described in Example 24.

Samples useful for analyses using the arrays of this invention include total RNA samples and mRNA samples. RNA samples can be prepared as described in Example 25. An RNA sample is reverse transcribed into cDNA and simultaneously labeled, i.e. with one member of a two-color fluorescent system, such as Cy3-dCTP/Cy5-dCTP as described in Example 26. The arrays are hybridized with the prepared sample and washed at appropriate

stringencies accounting for the choices of sample and probes of the array. The hybridization stringency can be higher when the probe sequence has higher homology with the gene it interrogates and when the probe is larger. A reference target, standard target, or other sample target for direct comparison may be prepared and hybridized simultaneously to the same array. A prepared sample will not degrade during hybridization and is labeled. Prepared samples are reverse transcribed and fluorescently labeled.

Hybridization results can be measured and analyzed using equipment and software available in the art as described in Example 27. Before finalizing data, preliminary results are preferably normalized by methods known in the art. An example of normalization appears in Example 29. Analysis includes determination of statistical significance. Measurement may include normalization and analysis, including statistical analysis. Resulting data are typically stored in computer files.

Mitochondrial biology expression microarrays are useful for detecting alterations in gene expression caused by alterations in mitochondrial biology. Although commercially available total genome expression arrays from companies such as Incyte Pharmaceuticals or Affymetrix contain probes for ten to twenty times as many genes as the arrays of this invention, the commercially available arrays have limitations. Several genes and probes that have been included on the arrays of this invention are not available on the commercial arrays. The commercial arrays are also very expensive and the large data sets resulting from them can be rather cumbersome to analyze and manipulate. The smaller, more focused arrays of this invention allow the expression patterns of hundreds of mitochondrial genes to be monitored quickly and efficiently. This study shows that a custom-designed microarray for mitochondrial biology expression studies, including probes for nuclear as well as mitochondrial genes, is an effective tool for the analysis of gene expression changes caused by alterations in functions resulting from a mutation in a gene involved in mitochondrial biology or other changes in metabolic state. The cell line experiments in Examples 13-17 and 20 have been particularly informative in demonstrating the specificity and sensitivity of the arrays of this invention while the mouse tissue experiments in Examples 8-12 and 16-17 have shown the consistency of the arrays of this invention.

Clones used to generate probes are listed in Tables 3-5. Clones range from about 1 kb to about 4 kb. The inserts of most clones have been sequenced on the 5' and 3' ends.

Sequences of the 5' and 3' ends of the clones are usually about 200 nt to about 800 nt and are provided herein. Probes may be generated via several methods. For example, the clones listed in Tables 3-5 may be obtained commercially, the inserts purified and used as probes. Alternatively, a 5' or 3' sequence given in the sequence listings hereof may be used to design an oligonucleotide which may be synthesized and used to probe a library to identify a cDNA or genomic clone that is equivalent to the clone used to generate the original sequence. This newly identified cDNA or genomic equivalent clone may be used to generate a probe. Alternatively, a pair of sequences from the sequence listings, representing the 5' and 3' ends of one clone, may be used to design PCR primers, which may be used to PCR amplify an isolated nucleic acid that is quivalent to the insert of the corresponding clone from which the 5' and 3' were derived. This isolated nucleic acid may be used as a probe. Probes should not contain a vector sequence that hybridizes with any sequence in a sample. Methods for designing PCR primers and designing oligonucleotides for screening libraries are known in the art.

EXAMPLES

Example 1

Human Mitochondrial Biology Array

A human mitochondrial biology array is made from clones representing 650 expressed sequences involved in mitochondrial biology. The clones used to make probes that are placed on the array are shown in Table 3 which references SEQ ID NOS:1-994 provided herein setting forth the 5' and 3' sequences from these clones. The clones identified in Table 3 are used to make a set of probes called Human Probe Set #1. Control sequences are also placed this array. Controls include, but are not limited to blanks, DMSO, probes derived from plant sequences, sequence(s) not involved in mitochondrial biology, and poly adenine (40-60 nucleotides long).

Table 3

SEQ ID NO	ResGen Clone ID	Gene	Complete Gene Name	GenBank	UniGene Cluster ID
1		mtDNA - 12S ribosome	Mitochondrial DNA 12S		
2		mtDNA - 16S ribosome	Mitochondrial DNA 16s		
3		mtDNA - ATP6	Mitochondrial DNA ATP6		
4		mtDNA - ATP8	Mitochondrial DNA ATP8		

5	i	mtDNA - COX1	Mitochondrial DNA COX1	į	
6		mtDNA - COX2	Mitochondrial DNA COX2		
7		mtDNA - COX3	Mitochondrial DNA COX3		
8		mtDNA - CYT B	Mitochondrial DNA CYTb	_	
9		mtDNA - ND1	Mitochondrial DNA ND1		
10		mtDNA - ND2	Mitochondrial DNA ND2		
11		mtDNA - ND3	Mitochondrial DNA ND3		
12		mtDNA - ND4	Mitochondrial DNA ND4		
13		mtDNA - ND4L	Mitochondrial DNA ND4L		•
14		mtDNA - ND5	Mitochondrial DNA ND5		
15		mtDNA - ND6	Mitochondrial DNA ND6	.	
16	213890	DECR1	2,4-dienoyl CoA reductase	H72937	Hs.81548
17	213890	DECR1	2,4-dienoyl CoA reductase	H72938	Hs.81548
18	588911	OAS1	2',5'-oligoadenylate synthetase 1	AA146773	Hs.82396
19	588911	OAS1	2',5'-oligoadenylate synthetase 1	AA146772	Hs.82396
20	1576254	2',5'-oligoadenylate synthetase 1	2',5'-oligoadenylate synthetase 1	AA954880	Hs.82396
21	1057786	OAS2 (splice-variant)	2'-5'oligoadenylate synthetase 2	AA568217	
22	2190112	A38234	2-OXOGLUTARATE DEHYDROGENASE E1 COMPONENT	AI610813	
23	814444	CRSP9	33 kDa transcriptional co-activator	AA459244	Hs.7558
24	814444	CRSP9	33 kDa transcriptional co-activator	AA459465	Hs.7558
25	896949	HMGCR	3-hydroxy-3-methylglutaryl-Coenzyme A reductase	AA779417	
26	290111	HMGCS1	3-hydroxy-3-methylglutaryl-Coenzyme A synthase 1 (soluble)	N62195	Hs.77910
27	290111	HMGCS1	3-hydroxy-3-methylglutaryl-Coenzyme A synthase 1 (soluble)	N76492	Hs.77910
28	757222	HMGCS2	3-hydroxy-3-methylglutaryl-Coenzyme A synthase 2 (mitochondrial)	AA496148	
29	757222	HMGCS2	3-hydroxy-3-methylglutaryl-Coenzyme A synthase 2 (mitochondrial)	AA496149	
30	109310	НААО	3-hydroxyanthranilate 3,4-dioxygenase	T80846	
31	109310	HAAO	3-hydroxyanthranilate 3,4-dioxygenase	T80921	
32	1635163	HAAO	3-hydroxyanthranilate 3,4-dioxygenase	AI005031	
33	66564	BDH	3-hydroxybutyrate dehydrogenase (heart, mitochondrial)	T67057	Hs.76893
34	66564	BDH	3-hydroxybutyrate dehydrogenase (heart, mitochondrial)	T67058	Hs.76893
35	838366	HMGCL	3-hydroxymethyl-3-methylglutaryl- Coenzyme A lyase (hydroxymethylglutaricaciduria)	AA458779	Hs.831
			3-hydroxymethyl-3-methylglutaryl- Coenzyme A lyase		
36	838366	HMGCL	(hydroxymethylglutaricaciduria)	AA458172	Hs.831
37	28469	OXCT	3-oxoacid CoA transferase	R13381	
38	28469	OXCT	3-oxoacid CoA transferase	R40897	
39	591540	PHGDH	3-phosphoglycerate dehydrogenase	AA158735	ļ
40	591540	PHGDH	3-phosphoglycerate dehydrogenase	AA159852	
41	266720	PDPKi	3-phosphoinositide dependent protein kinase-1	N22904	

42	266720	PDPK1	3-phosphoinositide dependent protein kinase-1	N31292	
43	666169	MTR	5-methyltetrahydrofolate-homocysteine methyltransferase	AA233640	
44	666169	MTR	5-methyltetrahydrofolate-homocysteine methyltransferase	AA233650	
45	814765	AKAP1	A kinase anchor protein, 149kD	AA454947	Hs.78921
46	814765	AKAP1	A kinase anchor protein, 149kD	AA455326	Hs.78921
47	2364633	A32422	A32422	AI744652	
48	2308263	A40487	A40487	AI671604	
49	2240514	AAP1'	AAP1'	AI637909	
50	2266774	ABF2	ABF2	AI590841	
51	36393	ACAT2	acetyl-Coenzyme A acetyltransferase 2 (acetoacetyl Coenzyme A thiolase)	R25823	·
			acetyl-Coenzyme A acetyltransferase 2		
52	36393	ACAT2	(acetoacetyl Coenzyme A thiolase)	R46821	į
53	45376	ACAA2	acetyl-Coenzyme A acyltransferase 2 (mitochondrial 3-oxoacyl-Coenzyme A thiolase)	H07926	Hs.32500
			acetyl-Coenzyme A acyltransferase 2	1107520	115.52500
54	45376	ACAA2	(mitochondrial 3-oxoacyl-Coenzyme A thiolase)	H08029	Hs.32500
55	262932	ACO2	Aconitase 2, mitochondrial	H99699	Hs.75900
56	366511	No Data	Actin, alpha 1, skeletal muscle	AA026609	
57	366511	No Data	Actin, alpha 1, skeletal muscle	AA026720	
58	867606	ACTB	actin, beta	AA780815	
59	428215	ACTL6	actin-like 6	AA001745	Hs. 103180
60	428215	ACTL6	actin-like 6	AA001815	Hs. 103180
61	896962	ACADS	acyl-Coenzyme A dehydrogenase, C-2 to C-3 short chain	AA676663	Hs.127610
62	298155	ACADM	Acyl-Coenzyme A dehydrogenase, C-4 to C-12 straight chain	N70794	Hs.79158
63	140131	ACADL	acyl-Coenzyme A dehydrogenase, long chain	R66005	Hs.1209
64	140131	ACADL	acyl-Coenzyme A dehydrogenase, long chain	R66006	Hs.1209
65	243100	ACADSB	acyl-Coenzyme A dehydrogenase, short/branched chain	H95792	Hs.81934
66	243100	ACADSB	acyl-Coenzyme A dehydrogenase, short/branched chain	H96140	Hs.81934
67	810358	ACADVL	Acyl-Coenzyme A dehydrogenase, very long chain	AA464163	Hs.82208
68	810358	ACADVL	Acyl-Coenzyme A dehydrogenase, very long chain	AA464228	Hs.82208
69	85450	ACOX2	acyl-Coenzyme A oxidase 2, branched chain	T71713	Hs.9795
70	85450	ACOX2	acyl-Coenzyme A oxidase 2, branched chain	T71782	Hs.9795
71	772304	ANT2	Adenine nucleotide translocator 2 (fibroblast)	AA404486	Hs.79172

		· ·		ı	1
١	ì		Adenine nucleotide translocator 2		TY. 50150
72	772304	ANT2	(fibroblast)	AA405477	Hs.79172
73	755855	ANT3	adenine nucleotide translocator 3 (liver)	AA496376	Hs.164280
74	755855	ANT3	adenine nucleotide translocator 3 (liver)	AA496654	Hs.164280
75	853570	ANT3	Adenine nucleotide translocator 3 (liver)	AA663439	Hs.164280
76	868757	AK1	Adenylate kinase 1	AA775325	Hs.76240
77	2010933	KAD2 HUMAN	ADENYLATE KINASE ISOENZYME 2	AI361029	
			ADP,ATP CARRIER PROTEIN, HEART/SKELETAL MUSCLE ISOFORM T1	R53337	Hs.2043
78	40026	ANT1		103337	113.2043
79	40026	ANT1	ADP,ATP CARRIER PROTEIN, HEART/SKELETAL MUSCLE ISOFORM T1	R53942	Hs.2043
80	46248	ADPRT	ADP-ribosyltransferase (NAD+; poly (ADP-ribose) polymerase)	H09923	Hs.177766
81	46248	ADPRT	ADP-ribosyltransferase (NAD+; poly (ADP-ribose) polymerase)	H09924	Hs.177766
82	589276	AFG3L2	AFG3 (ATPase family gene 3, yeast)-like 2	AA147320	Hs.29385
83	589276	AFG3L2	AFG3 (ATPase family gene 3, yeast)-like 2	AA147413	Hs.29385
				A 4 570 400	
84	1018253	AIF	AIF	AA570483	TT- 7(200
85	855624	ALDH1	aldehyde dehydrogenase 1, soluble	AA664101	Hs.76392
86	47853	ALDH4	aldehyde dehydrogenase 4 (glutamate gamma-semialdehyde dehydrogenase; pyrroline-5-carboxylate dehydrogenase)	H11346	Hs.77448
87	47853	ALDH4	aldehyde dehydrogenase 4 (glutamate gamma-semialdehyde dehydrogenase; pyrroline-5-carboxylate dehydrogenase)	H11369	Hs.77448
	197657	ALDH5	ALDEHYDE DEHYDROGENASE, MITOCHONDRIAL X PRECURSOR	R93550	Hs.169517
88		ALDH5	ALDEHYDE DEHYDROGENASE, MITOCHONDRIAL X PRECURSOR	R93551	Hs.169517
89	197657 1917741	AOX1	aldehyde oxidase 1	AI343711	Hs.81047
90	2154324	AGPS	alkylglycerone phosphate synthase	AI445035	
92	1520618	GABT HUMAN	AMINOBUTYRATE AMINOTRANSFERASE	AA910669	
93	813651	ALAS1	aminolevulinate, delta-, synthase 1	AA447761	Hs.78712
94		ALAS1	aminolevulinate, delta-, synthase 1	AA453691	Hs.78712
95	753346	ALAS2	aminolevulinate, delta-, synthase 2 (sideroblastic/hypochromic anemia)	AA406485	Hs.79103
96		ALAS2	aminolevulinate, delta-, synthase 2 (sideroblastic/hypochromic anemia)	AA410346	Hs.79103
97	248631	AMT	aminomethyltransferase (glycine cleavag system protein T)	N59532	Hs.102
98		AMT	aminomethyltransferase (glycine cleavag system protein T)	e N78273	Hs.102
99		ANT3	ANT3	AA916851	
100		API5L1	API5-like 1	AA035435	
100		API5L1	API5-like 1	AA035436	
102		API2	apoptosis inhibitor 2	R07870	Hs.127799
10		API2	apoptosis inhibitor 2	R07927	Hs.127799

ابمد		API3	apoptosis inhibitor 3	AI628066	
104	2285739	ARAL-1	ARAL-1	AA535370	
105	927606	XNHUDM	ASPARTATE AMINOTRANSFERASE	AI096615	
106	1704180	Alliebia	Ataxia telangiectasia mutated (includes		1
107	360778	ATM	complementation groups A, C and D)	AA016254	Hs.194382
10/	300778		Ataxia telangiectasia mutated (includes		
108	360778	ATM	complementation groups A, C and D)	AA016988	Hs.194382
100	300770		ATP SYNTHASE GAMMA CHAIN,		
109	845519	ATP5C1	MITOCHONDRIAL PRECURSOR	AA644234	Hs.155433
	0.000		ATP synthase, H+ transporting,		
			mitochondrial F0 complex, subunit b,	1 1 150555	77- 101101
110	813712	ATP5F1	isoform 1	AA453765	Hs.181101
			ATP synthase, H+ transporting,		[
1			mitochondrial F0 complex, subunit b, isoform 1	AA453849	Hs.181101
111	813712	ATP5F1		AA433049	113.101101
			ATP synthase, H+ transporting,		
		A TENS CO	mitochondrial F0 complex, subunit c (subunit 9) isoform 3	H47080	Hs.429
112	193106	ATP5G3			
1		(ATP synthase, H+ transporting, mitochondrial F0 complex, subunit c		
112	102106	ATP5G3	(subunit 9) isoform 3	H47164	Hs.429
113	193106	<u> </u>	ATP synthase, H+ transporting,		
		Ì	mitochondrial F0 complex, subunit c		
114	487373	ATP5G1	(subunit 9), isoform 1	AA046701	Hs.80986
11.4	107373		ATP synthase, H+ transporting,		
			mitochondrial F0 complex, subunit c		
115	487373	ATP5G1	(subunit 9), isoform 1	AA046489	Hs.80986
			ATP synthase, H+ transporting,		
			mitochondrial F0 complex, subunit c	AA455126	Hs.89399
116	809876	ATP5G2	(subunit 9), isoform 2	AA433120	118.09399
	1		ATP synthase, H+ transporting, mitochondrial F0 complex, subunit c	1	1
		ATP5G2	(subunit 9), isoform 2	AA464312	Hs.89399
117	809876	AIF3G2	ATP synthase, H+ transporting,		
110	825312	ATP5J	mitochondrial F0 complex, subunit F6	AA504540	Hs.73851
118	023312		ATP synthase, H+ transporting,		
111	9 825312	ATP5J	mitochondrial F0 complex, subunit F6	AA504465	Hs.73851
1.1	023312		ATP synthase, H+ transporting,		
-	-		mitochondrial F1 complex, beta	AA708298	Hs.25
12	0 392622	ATP5B	polypeptide	AA/08298	113.23
		, am an	ATP synthase, H+ transporting, mitochondrial F1 complex, delta subuni	AA669314	Hs.89761
12	1 856650	ATP5D	ATP synthase, H+ transporting,	121007514	120.05.02
			mitochondrial F1 complex, O subunit	1	
			(oligomycin sensitivity conferring		
12	1472150	ATP5O	protein)	AA873577	Hs.76572
12			ATP2B2	AI421603	Hs.89512
12		ATP50	ATP50	AI184610	
				110500	
1:	25 1173869	ATP5A1	ATP5A1	AA640573	-
			ATTRECT	AA507388	
	26 964121		ATP5C1	AI126623	
1	27 1736058	ATP5F1	ATP5F1	A120023	

128	307873	ATP2B4	ATPase, Ca++ transporting, plasma membrane 4	N93024	Hs.995
129	307873	ATP2B4	ATPase, Ca++ transporting, plasma membrane 4	W21376	Hs.995
130	1435103	ATP2A3	ATPase, Ca++ transporting, ubiquitous	AA857542	Hs.5541
131	266312	АТР7В	ATPase, Cu++ transporting, beta polypeptide (Wilson disease)	N26536	Hs.84999
132	266312	АТР7В	ATPase, Cu++ transporting, beta polypeptide (Wilson disease)	N35647	Hs.84999
133	384078	ATP6DV	ATPase, H+ transporting, lysosomal (vacuolar proton pump) 31kD	AA702541	Hs.106876
134	825170	ATP6A1	ATPase, H+ transporting, lysosomal (vacuolar proton pump), alpha polypeptide, 70kD, isoform 1	AA504160	Hs.255352
135	825170	ATP6A1	ATPase, H+ transporting, lysosomal (vacuolar proton pump), alpha polypeptide, 70kD, isoform 1	AA504159	Hs.255352
136	1323203	ATP6B2	ATPase, H+ transporting, lysosomal (vacuolar proton pump), beta polypeptide, 56/58kD, isoform 2	AA877194	Hs.1697
137	461522	ABCB7	ATP-binding cassette, sub-family B (MDR/TAP), member 7	AA705237	Hs.125856
138	1709773	MCX1	ATP-DEPENDENT CLP PROTEASE	AI131257	
139	1467799	KIAA0705	atrophin-1 interacting protein 1	AA883236	Hs.22599
140	852273	AZF1	AZF1	AA772863	
141	2367249	B42665	B42665	AI741963	
142	194384	BTF3	Basic transcription factor 3	R83000	Hs.101025
143	194384	BTF3	Basic transcription factor 3	R82957	Hs.101025
144	342181	BCL2	B-cell CLL/lymphoma 2	W63749	Hs.79241
145	342181	BCL2	B-cell CLL/lymphoma 2	W61100	Hs.79241
146	826182	BCL6	B-cell CLL/lymphoma 6 (zinc finger protein 51)	AA521434	
147	814899	BNIP3L	BCL2/adenovirus E1B 19kD-interacting protein 3-like	AA465697	Hs.132955
148	235938	BAK1	BCL2-antagonist/killer 1	H52672	
149	235938	BAK1	BCL2-antagonist/killer 1	H52673	
150	2125819	BAX	BCL2-associated X protein	AI565203	
151	1916575	BIK	BCL2-interacting killer (apoptosis- inducing)	AI347538	
152	1568561	BCL2L1	BCL2-like 1	AA931820	Hs.180372
153		BCS1	BCS1	AI670836	
154		BZRP	benzodiazapine receptor (peripheral)	AA455945	Hs.202
155	813444	BZRP	benzodiazapine receptor (peripheral)	AA455554	Hs.202
156	627125	BID	BH3 interacting domain death agonist	AA190401	
157	627125	BID	BH3 interacting domain death agonist	AA190546	
158	1573108	BCKDK	branched chain alpha-ketoacid dehydrogenase kinase	AA970731	Hs.20644
159	756490	BCAT2	branched chain aminotransferase 2, mitochondrial	AA436410	Hs.101408
160		BCAT2	branched chain aminotransferase 2, mitochondrial	AA481353	Hs.101408

161	740801	BCKDHA	branched chain keto acid dehydrogenase E1, alpha polypeptide (maple syrup urine disease)	AA477298	Hs.78950
			branched chain keto acid dehydrogenase E1, alpha polypeptide (maple syrup urine		
162	740801	BCKDHA	disease)	AA477297	Hs.78950
163	770835	всконв	Branched chain keto acid dehydrogenase E1, beta polypeptide (maple syrup urine disease)	AA427739	Hs.1265
164	770835	ВСКДНВ	Branched chain keto acid dehydrogenase E1, beta polypeptide (maple syrup urine disease)	AA434304	Hs.1265
165	129431	EST	BRCA2(?)	R11316	Hs.188591
166	129431	EST	BRCA2(?)	R11315	Hs.188591
167	83605	CPS1	carbamoyl-phosphate synthetase 1, mitochondrial	T61078	Hs.50966
			carbamoyl-phosphate synthetase 1,		
168	83605	CPS1	mitochondrial	T61180	Hs.50966
169	1675950	CRHU5	CARBONIC ANHYDRASE V PRECURSOR	AI052226	
170	744417	CRAT	Carnitine acetyltransferase	AA621218	Hs.12068
171	133565	CPT1A	carnitine palmitoyltransferase I, liver	R28631	Hs.29331
172	133565	CPT1A	carnitine palmitoyltransferase I, liver	R32561	Hs.29331
173	415978	CPT1A	carnitine palmitoyltransferase I, liver	W85710	Hs.29331
174	415978	CPT1A	carnitine palmitoyltransferase I, liver	W86378	Hs.29331
175	120106	CASP1	caspase 1, apoptosis-related cysteine protease (interleukin 1, beta, convertase)	T95052	
176	120106	CASP1	caspase 1, apoptosis-related cysteine protease (interleukin 1, beta, convertase)	T95149	
177	30170	CASP3	caspase 3, apoptosis-related cysteine protease	R14760	
178	30170	CASP3	caspase 3, apoptosis-related cysteine protease	R42530	
179	429574	CASP3	caspase 3, apoptosis-related cysteine protease	AA011445	
180	429574	CASP3	caspase 3, apoptosis-related cysteine protease	AA011446	
181	745143	CASP6	caspase 6, apoptosis-related cysteine protease	AA626710	
182	279691	CD2AP	CD2-associated protein	N48329	Hs.30490
183	279691	CD2AP	CD2-associated protein	N49054	Hs.30490
184	1558965	CDC2L	cholinesterase-related cell division controller	AA917769	Hs.155266
185	324885	C11ORF4	chromosome 11 open reading frame 4	W48701	Hs.75859
186	324885	C11ORF4	chromosome 11 open reading frame 4	W48700	Hs.75859
187	1573778	Cl1ORF13	chromosome 11 open reading frame 13	AA970526	Hs.72925
188	110772	C14ORF2	chromosome 14 open reading frame 2	T90621	Hs.109052
189	110772	C14ORF2	chromosome 14 open reading frame 2	T83147	Hs.109052
190	897448	C2ORF1	chromosome 2 open reading frame 1	AA489478	Hs.14454
191	1060841	CIT1	CITRATE SYNTHASE, MITOCHONDRIAL PRECURSOR	AA568724	
192		F10	COAGULATION FACTOR X PRECURSOR	N98524	Hs.47913

193	310519	F10	COAGULATION FACTOR X PRECURSOR	W31088	Hs.47913
193	310319		coproporphyrinogen oxidase		
194	436062	CPO	(coproporphyria, harderoporphyria)	AA700808	Hs.89866
195	161476	COQ5	COQ5	H25602	
196	161476	COQ5	COQ5	H25556	
197	2338136	COX11.1-11.2	COX11.1-11.2	AI703310	
198	1902314	COX15.1	COX15.1	AI301929	
199	1318021	COX5A	COX5A	AA769095	
200	2326019	COX5B	COX5B	AI688757	Hs.1342
200	2320019				
201	2301230	COX5B	COX5B	AI699318	
201	2097122	COX6A1	COX6A1	AI421088	
	986164	COX6A2	COX6A2	AA548887	
203	2277616	COX6B	COX6B	AI690478	
204	971851	COX7A1	COX7A1	AA515958	
205	937944	COX7A2	COX7A2	AA563616	
206	93/944	0011112	creatine kinase, mitochondrial 1		
000	2019469	CKMT1	(ubiquitous)	AI369378	Hs.153998
207	2019409	ORMITT	Creatine kinase, mitochondrial 2		
200	795965	CKMT2	(sarcomeric)	AA460480	Hs.80691
208	193903	- CILITA	Creatine kinase, mitochondrial 2		
200	795965	CKMT2	(sarcomeric)	AA461048	Hs.80691
209	771327	NIFS	cysteine desulfurase	AA476245	Hs.194692
210	771327	NIFS	cysteine desulfurase	AA476244	Hs.194692
211	196189	CYB5	Cytochrome b-5	R92281	Hs.83834
212 213	196189	CYB5	Cytochrome b-5	R91950	Hs.83834
213	190109	0120	CYTOCHROME C OXIDASE		
1			POLYPEPTIDE VIA-LIVER		400-44
214	840894	COX6A1	PRECURSOR	AA482243	Hs.180714
			CYTOCHROME C OXIDASE		ļ
		0077614	POLYPEPTIDE VIA-LIVER PRECURSOR	AA482340	Hs.180714
215	840894	COX6A1	cytochrome c oxidase subunit Va	AA490735	110.100711
216		COX5A	cytochrome c oxidase subunit Va	AA491224	
217		COX5A	Cytochrome c oxidase subunit VIb	N71160	Hs.174031
218	298965	COX6B	Cytochrome c oxidase subunit VIb	W05541	Hs.174031
219		COX6B	cytochrome c oxidase subunit VIb	AA872391	Hs.174031
220		COX6B	cytochrome c oxidase subunit VID	AA456931	Hs.74649
221		COX6C	eytochrome c oxidase subunit VIC	AA457006	Hs.74649
222	838568	COX6C		AA437000	113.74042
İ		gover 1	cytochrome c oxidase subunit VIIa polypeptide 1 (muscle)	AA872125	Hs.114346
223	1475803	COX7A1		PHIOTETES	115.2 1 15 10
		00377.10	cytochrome c oxidase subunit VIIa polypeptide 2 (liver)	AI002403	Hs.182684
224		COX7A2	cytochrome c oxidase subunit VIIb	AA629999	Hs.75752
22:		COX7B	cytochrome c oxidase subunit VIIc	AA629719	Hs.3462
22		COX7C	cytochrome c oxidase subunit VIII	AA862813	Hs.81097
22	7 1469230	COX8		111002013	1
			Cytochrome c oxidase subunit X (heme A: farnesyltransferase	H21868	Hs.77513
228	160126	COX10		1121000	1
			Cytochrome c oxidase subunit X (heme	H21869	Hs.77513
22		COX10	A: farnesyltransferase cytochrome c-1	AA865265	Hs.697
23		CYC1	Cytochrome P450 IIIA7 (P450-HFLa)	R91078	Hs.172323
23	1 194949	CYP3A7	Cytochrome P430 IIIA/ (P430-HFLa)	1010/3	110.1/2020

232	194949	CYP3A7	Cytochrome P450 IIIA7 (P450-HFLa)	R91077	Hs.172323
233	1724630	S14367	CYTOCHROME P450 XIA1	AI183397	
234	85561	CYP2A7	cytochrome P450, subfamily IIA (phenobarbital-inducible), polypeptide 7	T73031	Hs.252937
235	85561	CYP2A7	cytochrome P450, subfamily IIA (phenobarbital-inducible), polypeptide 7	T72259	Hs.252937
236	246619	CYP2C8	Cytochrome P450, subfamily IIC (mephenytoin 4-hydroxylase)	N53136	
237	246619	CYP2C8	Cytochrome P450, subfamily IIC (mephenytoin 4-hydroxylase)	N58566	
238	195712	CYP2C9	Cytochrome P450, subfamily IIC (mephenytoin 4-hydroxylase), polypeptide 9	R89492	Hs.167529
239	195712	CYP2C9	Cytochrome P450, subfamily IIC (mephenytoin 4-hydroxylase), polypeptide 9	R89491	Hs.167529
240	1467195	CYP11B1	cytochrome P450, subfamily XIB (steroid 11-beta-hydroxylase), polypeptide 1	AA884709	Hs.2610
241	149737	EST	Cytochrome P450, subfamily XIX (aromatization of androgens)	H00592	Hs.141142
242	149737	EST	Cytochrome P450, subfamily XIX (aromatization of androgens)	R82738	Hs.141142
243	284620	EST	Cytochrome P450, subfamily XXI (steroid 21-hydroxylase, congenital adrenal hyperplasia)	N64794	Hs.124918
244	284620	EST	Cytochrome P450, subfamily XXI (steroid 21-hydroxylase, congenital adrenal hyperplasia)	N77388	Hs.124918
245	266146	CYP24	cytochrome P450, subfamily XXIV (vitamin D 24-hydroxylase)	N21576	Hs.89663
246	266146	CYP24	cytochrome P450, subfamily XXIV (vitamin D 24-hydroxylase)	N30976	Hs.89663
247	295843	CYP27A1	cytochrome P450, subfamily XXVIIA (steroid 27-hydroxylase, cerebrotendinous xanthomatosis), polypeptide 1	N66957	Hs.82568
248	2043415	DAPK1	death-associated protein kinase 1	AI371096	Hs.153924
249	2364396	DEHUH2	DEHUH2	AI740677	
250	125722	DGUOK	deoxyguanosine kinase	R07560	Hs.77494
251	125722 2096376	DGUOK. S25665	deoxyguanosine kinase DIHYDROLIPOAMIDE ACETYLTRANSFERASE COMPONENT	R07506 AI419467	Hs.77494
253	813648	DLD	dihydrolipoamide dehydrogenase (E3 component of pyruvate dehydrogenase complex, 2-oxo-glutarate complex, branched chain keto acid dehydrogenase complex)	AA453679	Hs.74635
254	813648	DLD	dihydrolipoamide dehydrogenase (E3 component of pyruvate dehydrogenase complex, 2-oxo-glutarate complex, branched chain keto acid dehydrogenase complex)	AA447748	Hs.74635

			1		ì
			dihydrolipoamide S-acetyltransferase (E2		
			component of pyruvate dehydrogenase		
255	271006_	DLAT	complex)	N29901	Hs.115285
\Box			dihydrolipoamide S-acetyltransferase (E2		
			component of pyruvate dehydrogenase		
256	271006	DLAT	complex)	N42953	Hs.115285
			Dihydrolipoamide S-succinyltransferase		
			(E2 component of 2-oxo-glutarate		
257	815564	DLST	complex)	AA456824	Hs.196416
			Dihydrolipoamide S-succinyltransferase		
			(E2 component of 2-oxo-glutarate		
258	815564	DLST	complex)	AA456899	Hs.196416
i i		DIHYDROLIPOAMIDE			
1 1		SUCCINYLTRANSFERAS	DIHYDROLIPOAMIDE		
259	1308945	Е	SUCCINYLTRANSFERASE	AA748401	
		DIHYDROLIPOAMIDE			
.		SUCCINYLTRANSFERAS	DİHYDROLIPOAMIDE		
260	1308945	E	SUCCINYLTRANSFERASE	AA746285	
200	1300543		BOCCH I BILL HIST BILL BE	71117-10205	
001	417205	Dihydroorotate dehydrogenase	Dihydroorotate dehydrogenase	W88472	Hs.125846
261	417385		Diffydfoofolate deifydfogeliase	W00472	HS.123640
		Dihydroorotate	73	*********	**
262	417385	dehydrogenase	Dihydroorotate dehydrogenase	W89035	Hs.125846
		dihydroorotate			
263	611027	dehydrogenase	dihydroorotate dehydrogenase	AA173122	Hs.94925
1		dihydroorotate			
264	611027	dehydrogenase	dihydroorotate dehydrogenase	AA173225	Hs.94925
265	884539	DKFZP566D143	DKFZP566D143 protein	AA629804	
266	630013	MSH2	DNA repair protein MSH2	AA219060	Hs.78934
267	630013	MSH2	DNA repair protein MSH2	AA219061	Hs.78934
268	666425	PLCG1	DNA topoisomerase I	AA232856	Hs.317
269	666425	PLCG1	DNA topoisomerase I	AA233029	Hs.317
270	810787	HSPF1	DNAJ PROTEIN HOMOLOG 1	AA481758	Hs.82646
271	810787	HSPF1	DNAJ PROTEIN HOMOLOG 1	AA481022	Hs.82646
	010707		dysferlin, limb girdle muscular dystrophy		122,020,10
272	1914863	DYSF	2B (autosomal recessive)	AI310142	Hs.143897
2/2	1914605		<u> </u>	111310172	113.143097
			dystrophin (muscular dystrophy,		
1 1			Duchenne and Becker types), includes		
			DXS142, DXS164, DXS206, DXS230,		
	## #	DIA.	DXS239, DXS268, DXS269, DXS270,	A A 405C40	TT- 1.00.400
273	773399	DMD	DXS272	AA425649	Hs.169470
			dystrophin (muscular dystrophy,		
			Duchenne and Becker types), includes		
			DXS142, DXS164, DXS206, DXS230,		
			DXS239, DXS268, DXS269, DXS270,		
274	773399	DMD	DXS272	AA427831	Hs.169470
			Dystrophin (muscular dystrophy,		
		1	Duchenne and Becker types), includes		
			DXS142, DXS164, DXS206, DXS230,		
		1	DXS239, DXS268, DXS269, DXS270,		
275	796197	DMD	DXS272	AA461118	Hs.169470
ئنت					·

		D) (D	Dystrophin (muscular dystrophy, Duchenne and Becker types), includes DXS142, DXS164, DXS206, DXS230, DXS239, DXS268, DXS269, DXS270,	AA461435	Hs.169470
276	796197	DMD	DXS272 early growth response 2 (Krox-20	AA401455	П8.109470
277	781017	EGR2	(Drosophila) homolog)	AA446027	Hs.1395
	#0101 5	EGR2	early growth response 2 (Krox-20 (Drosophila) homolog)	AA446300	Hs.1395
278	781017		ectodermal-neural cortex (with BTB-like		
279	180512	ENC1	domain) ELECTRON TRANSFER	R85090	Hs.104925
280	1744035	A31998	FLAVOPROTEIN ALPHA-SUBUNIT	AI192719	
281	2274670	S32482	ELECTRON TRANSFER FLAVOPROTEIN BETA-SUBUNIT	AI683530	
282	2267229	S62767	ELONGATION FACTOR TU	A1609398	
	745542	ECHS1	enoyl Coenzyme A hydratase, short chain, 1, mitochondrial	AA626255	Hs.76394
283	344272	EMP3	epithelial membrane protein 3	W73810	Hs.9999
285	344272	EMP3	epithelial membrane protein 3	W73748	Hs.9999
286	253725	EST	EST	N21972	Hs.43052
287	287569	EST	EST	N62122	Hs.83313
288	287569	EST	EST	N78351	Hs.83313
289	489755	EST	EST	AA099554	Hs.246174
290		EST	EST	AA101991	Hs.246174
291	609989	No Data	EST	AA169176	
292	609989	No Data	EST	AA169296	
293	511012	AGPS	EST - putative alkylglycerone phosphate synthase	AA099787	
294	511012	AGPS	EST - putative alkylglycerone phosphate synthase	AA102257	
-			EST, Weakly similar to predicted using		
295	449504	EST	Genefinder [C.elegans]	AA777928	Hs.121993
296	47005	EST	ESTs	H09825	Hs.6818
297	47005	EST	ESTs	H09920	Hs.6818
298	79655	EST	ESTs	T62655	Hs.11039
299	79655	EST	ESTs	T62509	Hs.11039
300	126229	EST	ESTs	R06313	Hs.77677
301	126229	EST	ESTs	R06258	Hs.77677
302		EST	ESTs	R16545	Hs.70333
303		EST	ESTs	R16603	Hs.70333
304		EST	ESTs	R86713	Hs.87595
305		EST	ESTs	R86712	Hs.87595
306		EST	ESTs	N59553	Hs.8941
307		EST	ESTs ESTs	N78295 N22302	Hs.8941 Hs.177861
308		EST	ESTs ESTs	N75187	Hs.177861
309		LOC54675	ESTS	N29545	Hs.3569
310			ESTS	N67991	Hs.30487
311		EST	ESTS	N80413	Hs.30487
312		EST	ESTS	W61374	Hs.11317
314		EST EST	ESTS	AA034062	Hs.38750
141/	429942	1 E91	1 1015	TATAUJTUUZ	1 170.00/00

316	784214	EST	ESTs	AA446980	Hs.14018
317	898222	EST	ESTs	AA598602	Hs.13434
318	1636741	EŠTs	ESTs	AI017846	Hs.169539
		EST	ESTs, Highly similar to ARGINYL- TRNA SYNTHETASE, MITOCHONDRIAL PRECURSOR [Saccharomyces cerevisiae]	T83996	
319	113206		EST's, Highly similar to ARGINYL- TRNA SYNTHETASE, MITOCHONDRIAL PRECURSOR	183990	Hs.15395
320	113206	EST	[Saccharomyces cerevisiae] ESTs, Highly similar to ATP	T83997	Hs.15395
321	825386	ATP5JD	SYNTHASE D CHAIN, MITOCHONDRIAL [Bos taurus]	AA504246	Hs.64593
			ESTs, Highly similar to ATP SYNTHASE E CHAIN, MITOCHONDRIAL [Cricetulus		
322	782439	ATP5I	longicaudatus] ESTs, Highly similar to ATP SYNTHASE E CHAIN,	AA431433	Hs.85539
323	782439	ATP5I ATP5E	MITOCHONDRIAL [Cricetulus longicaudatus] ESTs, Highly similar to ATP	AA431836	Hs.85539
324	434968	AIFJE	SYNTHASE EPSILON CHAIN, MITOCHONDRIAL PRECURSOR [Bos taurus]	AA700688	Hs.177530
			ESTs, Highly similar to ATPASE INHIBITOR, MITOCHONDRIAL		
325	82874 82874	EST	PRECURSOR [Rattus norvegicus] ESTs, Highly similar to ATPASE INHIBITOR, MITOCHONDRIAL PRECURSOR [Rattus norvegicus]	T69273 T69348	Hs.241336 Hs.241336
327	290753	CS	ESTs, Highly similar to CITRATE SYNTHASE, MITOCHONDRIAL PRECURSOR [Sus scrofa]	N67639	Hs.239760
328	290753	CS	ESTs, Highly similar to CITRATE SYNTHASE, MITOCHONDRIAL PRECURSOR [Sus scrofa]	W01297	Hs.239760
329	731308	CS	ESTs, Highly similar to CITRATE SYNTHASE, MITOCHONDRIAL PRECURSOR [Sus scrofa]	AA416759	Hs.239760
330	731308	CS	ESTs, Highly similar to CITRATE SYNTHASE, MITOCHONDRIAL PRECURSOR [Sus scrofa]	AA416746	Hs.239760
331	283943	EST	ESTs, Highly similar to ELONGATION FACTOR G, MITOCHONDRIAL PRECURSOR [Rattus norvegicus]	N50802	Hs.41066
332	283943	EST	ESTs, Highly similar to ELONGATION FACTOR G, MITOCHONDRIAL PRECURSOR [Rattus norvegicus]	N55159	Hs.41066
333	359723	EST	ESTs, Highly similar to ELONGATION FACTOR G, MITOCHONDRIAL PRECURSOR [Rattus norvegicus]	AA011122	Hs.41066
334	359723	EST	ESTs, Highly similar to ELONGATION FACTOR G, MITOCHONDRIAL PRECURSOR [Rattus norvegicus]	AA010761	Hs.41066

			ESTs, Highly similar to HYPOTHETICAL 16.5 KD PROTEIN IN PAS8-EGT2 INTERGENIC REGION		XX. 2595
335	430733	EST	[Saccharomyces cerevisiae]	AA677960	Hs.3585
336	1114960	ETFDH	ESTs, Highly similar to HYPOTHETICAL 29.2 KD PROTEIN IN PHD1-PTM1 INTERGENIC REGION [Saccharomyces cerevisiae]	AA602015	Hs.30661
337	1030791	EST	ESTs, Highly similar to HYPOTHETICAL 44.9 KD PROTEIN IN ERG7-NMD2 INTERGENIC REGION [Saccharomyces cerevisiae]	AA609009	Hs.63304
	950700	PP	ESTs, Highly similar to INORGANIC PYROPHOSPHATASE [Bos taurus]	AA608572	Hs.184011
338		EST	ESTS, Highly similar to ISOLEUCYL- TRNA SYNTHETASE, MITOCHONDRIAL [Saccharomyces cerevisiae]	W37993	Hs.55609
339	322218	E31	ESTs, Highly similar to ISOLEUCYL- TRNA SYNTHETASE, MITOCHONDRIAL [Saccharomyces		4
340	322218	EST	cerevisiae]	W37992	Hs.55609
3.0			ESTs, Highly similar to MITOCHONDRIAL 60S RIBOSOMAL PROTEIN L2 PRECURSOR		
341	417801	LOC51264	[Saccharomyces cerevisiae]	W88752	Hs.7736
342	417801	LOC51264	ESTs, Highly similar to MITOCHONDRIAL 60S RIBOSOMAL PROTEIN L2 PRECURSOR [Saccharomyces cerevisiae]	W88848	Hs.7736
		EST	ESTs, Highly similar to MITOCHONDRIAL CARRIER PROTEIN RIM2 [Saccharomyces cerevisiae]	AA479944	Hs.42484
343	753602		ESTs, Highly similar to MITOCHONDRIAL CARRIER PROTEIN RIM2 [Saccharomyces	AA478847	Hs.42484
344	753602	EST	Cerevisiae] ESTs, Highly similar to MITOCHONDRIAL IMPORT INNER MEMBRANE TRANSLOCASE SUBUNIT TIM17 [Saccharomyces		
345	629916	TIM17B	cerevisiae] ESTs, Highly similar to MITOCHONDRIAL IMPORT INNER MEMBRANE TRANSLOCASE SUBUNIT TIM17 [Saccharomyces	AA219179	Hs.19105
340	629916	TIM17B	cerevisiae]	AA219178	Hs.19105
			ESTs, Highly similar to MITOCHONDRIAL RNA SPLICING PROTEIN MRS3 [Saccharomyces		
34	7 191826	EST	cerevisiae]	H40449	Hs.34401
			ESTs, Highly similar to MITOCHONDRIAL RNA SPLICING PROTEIN MRS3 [Saccharomyces	·	** 6.4405
34	8 191826	EST	cerevisiae]	H40448	Hs.34401

			ESTs, Highly similar to MITOCHONDRIAL RNA SPLICING PROTEIN MSR4 [Saccharomyces	W20000	TT- 24401
349	70201	EST	cerevisiae]	T50082	Hs.34401
			ESTs, Highly similar to MITOCHONDRIAL RNA SPLICING PROTEIN MSR4 [Saccharomyces	T50019	Hs.34401
350	70201	EST	cerevisiae] ESTs, Highly similar to	130019	113.5-1-01
351	488386	EST	MITOCHONDRIAL RNA SPLICING PROTEIN MSR4 [Saccharomyces cerevisiae]	AA046639	Hs.34401
331	400300		ESTs, Highly similar to MITOCHONDRIAL RNA SPLICING PROTEIN MSR4 [Saccharomyces		
352	488386	EST	, cerevisiae]	AA046778	Hs.34401
353	611467	NDUFS6	ESTs, Highly similar to MITOCHONDRIAL TRIFUNCTONAL ENZYME ALPHA SUBUNIT PRECURSOR [Homo sapiens] ESTs, Highly similar to	AA176453	Hs.49767
354	611467	NDUFS6	MITOCHONDRIAL TRIFUNCTONAL ENZYME ALPHA SUBUNIT PRECURSOR [Homo sapiens]	AA176931	Hs.49767
355	179336	SDHA	ESTs, Highly similar to SUCCINATE DEHYDROGENASE [Homo sapiens]	H50345	Hs.469
356	179336	SDHA	ESTs, Highly similar to SUCCINATE DEHYDROGENASE [Homo sapiens]	H50378	Hs.469
357	843335	SUCLG1	ESTs, Highly similar to SUCCINYL- COA LIGASE [Rattus norvegicus]	AA485965	Hs.7043
358	843335	SUCLG1	ESTs, Highly similar to SUCCINYL- COA LIGASE [Rattus norvegicus]	AA489575	Hs.7043
359	825229	LOC51017	ESTs, Highly similar to CGI-113 protein [H.sapiens]	AA504139	Hs.19077
360	825229	LOC51017	ESTs, Highly similar to CGI-113 protein [H.sapiens]	AA504401	Hs.19077
361	814271	Est	ESTs, Highly similar to CGI-116 protein [H.sapiens]	AA459002	Hs.18885
362	814271	Est	ESTs, Highly similar to CGI-116 protein [H.sapiens]	AA459222	Hs.18885
	811062	LOC51629	ESTs, Highly similar to CGI-69 protein [H.sapiens]	AA485441	Hs.237924
363	811062	LOC51629	ESTs, Highly similar to CGI-69 protein [H.sapiens]	AA485607	Hs.237924
365	417803	EST	ESTs, Highly similar to hypothetical protein [H.sapiens]	W88753	Hs.166406
366	417803	EST	ESTs, Highly similar to hypothetical protein [H.sapiens]	W88859	Hs.166406
367	878316	ESTs	ESTs, Highly similar to small zinc finger- like protein [H.sapiens]	AA670296	Hs.109571
307	0.0313		ESTs, Moderately similar to 3- OXOACYL-[ACYL-CARRIER- PROTEIN] SYNTHASE II [Escherichia		
368	504689	EST	coli]	AA149172	Hs.55781

		EST	ESTs, Moderately similar to 3- OXOACYL-[ACYL-CARRIER- PROTEIN] SYNTHASE II [Escherichia coli]	AA149171	Hs.55781
369	504689	EST	ESTs, Moderately similar to ABC1	111117171	113.55701
	1		PROTEIN PRECURSOR		
370	238907	D6S52E	[Saccharomyces cerevisiae]	H67876	Hs.243960
			ESTs, Moderately similar to ABC1		
		DCG50F	PROTEIN PRECURSOR [Saccharomyces cerevisiae]	H67202	Hs.243960
371	238907	D6S52E		1107202	113.2 13700
			ESTs, Moderately similar to 5- AMINOLEVULINIC ACID SYNTHASE		
			MITOCHONDRIAL PRECURSOR,		
372	435314	EST	ERYTHROID-SPECIFIC [H.sapiens]	AA699919	Hs.114018
			ESTs, Moderately similar to CGI-24	A A 497400	
373	839027	EST	protein [H.sapiens]	AA487499	
274	839027	EST	ESTs, Moderately similar to CGI-24 protein [H.sapiens]	AA487715	1
374	839021	251	ESTs, Moderately similar to MSG1-		,
375	321354	EST	related protein [H.sapiens]	W32403	Hs.18349
			ESTs, Moderately similar to MSG1-		
376	321354	EST	related protein [H.sapiens]	W32980	Hs.18349
377	824911	EST	ESTs, Weakly similar to /prediction	AA489022 AA489118	Hs.5080 Hs.5080
378	824911	EST	ESTs, Weakly similar to /prediction ESTs, Weakly similar to anon2A5	AA469116	HS.3060
379	85384	EST	[D.melanogaster]	T71965	Hs.10964
3/3	03307		ESTs, Weakly similar to anon2A5		
380	85384	EST	[D.melanogaster]	T72105	Hs.10964
		Tom	ESTs, Weakly similar to aralar1 [H.sapiens]	AA426113	Hs.183047
381	757265	FLJ20420	ESTs, Weakly similar to head-elevated	AA420113	113.163047
382	490753	12520.20	expression in 0.9 kb [D.melanogaster]	AA133166	Hs.6693
302	+50733		ESTs, Weakly similar to mitochondrial		
383	343555	EST	inner membrane protease 1 [S.cerevisiae]	W69379	Hs.62669
			ESTs, Weakly similar to mitochondrial	10/602/79	17-62660
384	343555	EST	inner membrane protease 1 [S.cerevisiae]	W69378	Hs.62669
1			ESTs, Weakly similar to MITOCHONDRIAL PRECURSOR	ļ	[
1			PROTEINS IMPORT RECEPTOR		
385	198312_	KIAA0719	[Neurospora crassa]	R94191	Hs.21198
			ESTs, Weakly similar to	ļ	
	1		MITOCHONDRIAL PRECURSOR	ļ	
200	100212	KIAA0719	PROTEINS IMPORT RECEPTOR [Neurospora crassa]	R94290	Hs.21198
386	198312	NIAA0/13	ESTs, Weakly similar to	10 (250	12312129
}	1		MITOCHONDRIAL PRECURSOR		
			PROTEINS IMPORT RECEPTOR		
387	511257	KIAA0719	[Neurospora crassa]	AA088722	Hs.21198
			ESTs, Weakly similar to		
			MITOCHONDRIAL PRECURSOR PROTEINS IMPORT RECEPTOR		
388	511257	KIAA0719	[Neurospora crassa]	AA088799	Hs.21198

		,	. 1	1	ı
389	324651	EST	ESTs, Weakly similar to MITOCHONDRIAL RESPIRATORY FUNCTION PROTEIN 1 [Saccharomyces cerevisiae]	W47099	Hs.19513
369		DOD.	ESTs, Weakly similar to MITOCHONDRIAL RESPIRATORY FUNCTION PROTEIN 1 [Saccharomyces cerevisiae]	W47223	Hs.19513
390	324651	EST	ESTs, Weakly similar to MSF1		-
391	429222	EST	PROTEIN [S.cerevisiae]	AA004210	Hs.3945
392	429222	EST	ESTs, Weakly similar to MSF1 PROTEIN [S.cerevisiae]	AA007411	Hs.3945
393	754537	EST	ESTs, Weakly similar to myosin beta heavy chain, cardiac and skeletal muscle [H.sapiens]	AA406291	
		EST	ESTs, Weakly similar to myosin beta heavy chain, cardiac and skeletal muscle [H.sapiens]	AA421979	
394	754537 897557	EST	ESTs, Weakly similar to myosin beta heavy chain, cardiac and skeletal muscle [H.sapiens]	AA497024	Hs.179817
396	897557	EST	ESTs, Weakly similar to myosin beta heavy chain, cardiac and skeletal muscle [H.sapiens]	AA489606	Hs.179817
397	812169	EST	ESTs, Weakly similar to PROBABLE MITOCHONDRIAL 40S RIBOSOMAL PROTEIN S5 [S.cerevisiae]	AA456042	Hs.7807
398	208531	EST	ESTs, Weakly similar to PUTATIVE MITOCHONDRIAL CARRIER YBR291C [Saccharomyces cerevisiae]	H61979	Hs.7994
399	208531	EST	ESTs, Weakly similar to PUTATIVE MITOCHONDRIAL CARRIER YBR291C [Saccharomyces cerevisiae]	H61978	Hs.7994
400	435509	tRNAleu	ESTs, Weakly similar to similar to leucyl-tRNA synthetase [C.elegans]	AA701379	
401	43662	EST	ESTs, Weakly similar to similar to mitochrondrial carrier protein [C.elegans]	H05645	Hs.21262
402		EST	ESTs, Weakly similar to similar to mitochrondrial carrier protein [C.elegans]	H05644	Hs.21262
		EST	ESTs, Weakly similar to Similar to NAD(P) transhydrogenase, mitochondrial [C.elegans]		Hs.106620
403		EST	ESTs, Weakly similar to Similar to NAD(P) transhydrogenase, mitochondria [C.elegans]	l R16232	Hs.106620
404		EST	ESTs, Weakly similar to VISC PROTEIN [Escherichia coli]	T74882	Hs.12239
405	84880		ESTs, Weakly similar to VISC PROTEIN		Hs.12239
406	84880	EST	[Escherichia coli] Fanconi anemia, complementation group		Hs.86297
40′		FANCA	A ferredoxin 1	AA187349	Hs.744
40			ferredoxin 1	AA188427	Hs.744
409			ferrochelatase (protoporphyria)	AA025142	Hs.26

411	365149	FECH	ferrochelatase (protoporphyria)	AA025157	Hs.26
412	1469138	FGA	Fibrinogen, A alpha polypeptide	AA865707	Hs.90765
413	131839	FOLR1	folate receptor 1 (adult)	R24635	Hs.73769
414	131839	FOLR1	folate receptor 1 (adult)	R24530	Hs.73769
415	146605	FPRL1	formyl peptide receptor-like 1	R80041	Hs.99855
416	146605	FPRL1	formyl peptide receptor-like 1	R79948	Hs.99855
417	490600	FOSL2	FOS-like antigen 2	AA101617	Hs.155210
418	490600	FOSL2	FOS-like antigen 2	AA101616	Hs.155210
419	279790	FMR1	Fragile X mental retardation 1	N48355	Hs.89764
420	279790	FMR1	Fragile X mental retardation 1	N49132	Hs.89764
421	669419	FRDA	Friedreich ataxia	AA253413	Hs.95998
422	669419	FRDA	Friedreich ataxia	AA253388	Hs.95998
423	469412	FH	Fumarate hydratase	AA026917	Hs.75653
424	469412	FH	Fumarate hydratase	AA026918	Hs.75653
			fusion, derived from t(12;16) malignant		
425	842839	FUS	liposarcoma	AA486284	
			fusion, derived from t(12;16) malignant		
426	842839	FUS	liposarcoma	AA489305	
			Gamma-actin, enteric smooth muscle		
427	81289	ACTG2	form	T60048	Hs.77443
			gap junction protein, beta 1, 32kD		
			(connexin 32, Charcot-Marie-Tooth	2750004	
428	288663	GJB1	neuropathy, X-linked)	N62394	
			gap junction protein, beta 1, 32kD		1
	000660	CTD1	(connexin 32, Charcot-Marie-Tooth	N79360	
429	288663	GJB1	neuropathy, X-linked) GCDH HUMAN	AI693352	
430	2338827	GCDH_HUMAN		A1095552	
431	66534	GCKR	glucokinase (hexokinase 4) regulatory	T67007	Hs.89771
1.52			glucokinase (hexokinase 4) regulatory		
432	66534	GCKR	protein	T67006	Hs.89771
			glucosamine (N-acetyl)-6-sulfatase		
433	471498	GNS	(Sanfilippo disease IIID)	AA035347	Hs.2703
434	122636	GGPC	Glucose-6-phosphatase	T98886	Hs.242
435	122636	GGPC	Glucose-6-phosphatase	T98887	Hs.242
436	166236	G6PD	glucose-6-phosphate dehydrogenase	R87497	
437	166236	G6PD	glucose-6-phosphate dehydrogenase	R88192	
438	753775	GMPR	Glucose-6-phosphate dehydrogenase	AA406242	Hs.1435
439	753775	GMPR	Glucose-6-phosphate dehydrogenase	AA410375	Hs.1435
440	361565	GLUD1	glutamate dehydrogenase 1	AA018372	Hs.77508
441	361565	GLUD1	glutamate dehydrogenase 1	AA017175	Hs.77508
1		507	glutamic-oxaloacetic transaminase 1,	H22856	Ha 507
442	51702	GOT1	soluble (aspartate aminotransferase 1)	H22856	Hs.597
	51500	COTI	glutamic-oxaloacetic transaminase 1, soluble (aspartate aminotransferase 1)	H22855	Hs.597
443	51702	GOT1		H22033	115.557
			Glutamic-oxaloacetic transaminase 2,		
	041000	COTT	mitochondrial (aspartate aminotransferase	AA487739	Hs.170197
444	841370	GOT2		AA-0/133	113.170197
	1		Glutamic-oxaloacetic transaminase 2,		
	941000	GOT2	mitochondrial (aspartate aminotransferase 2)	AA487521	Hs.170197
445		GPX1	glutathione peroxidase 1	AA485362	Hs.76686
446		GPX1 GPX1	glutathione peroxidase 1	AA485517	Hs.76686
447	810999	J GFAI	Brarantone beroxidase 1	1 222703317	, 225,70000

4401	587847	GPX2	Giutaunone perestrates 2, gas a	AA135289	Hs.2704
448		GPX2	Glutathione peroxidase 2, gastrointestinal	AA135152	Hs.2704
449	587847	GPX3		AA664180	Hs.172153
450	855523	GPX3	glutathione peroxidase 3 (plasma)	AI147534	Hs.172153
451	1555659	GSR		AA777289	Hs.121524
452	448619	GSS		AA463458	Hs.82327
453	811792	GPD1	glycerol-3-phosphate dehydrogenase 1 (soluble)	AA192547	Hs.255718
454	628418		glycerol-3-phosphate dehydrogenase 1 (soluble)	AA193116	Hs.255718
455	628418	GPD1	glycerol-3-phosphate dehydrogenase 2		
456	428756	GPD2	(mitochondrial)	AA005219	Hs.93201
457	428756	GPD2	glycerol-3-phosphate dehydrogenase 2 (mitochondrial)	AA005218	Hs.93201
458	42558	GATM	glycine amidinotransferase (L- arginine:glycine amidinotransferase)	R61229	Hs.75335
459	42558	GATM	glycine amidinotransferase (L- arginine:glycine amidinotransferase)	R61228	Hs.75335
460	134748	GCSH	glycine cleavage system protein H (aminomethyl carrier)	R28294	Hs.77631
461	134748	GCSH	glycine cleavage system protein H (aminomethyl carrier)	R28081	Hs.77631
462	248261	GLDC	glycine dehydrogenase (decarboxylating; glycine decarboxylase, glycine cleavage system protein P)	N78083	Hs.27
463	248261	GLDC	glycine dehydrogenase (decarboxylating; glycine decarboxylase, glycine cleavage system protein P)	N58494	Hs.27
464	45632	GYS1	glycogen synthase 1 (muscle)	H08446	Hs.772
465	45632	GYS1	- glycogen synthase 1 (muscle)	H08732	Hs.772
466	245920	GYS2	glycogen synthase 2 (liver)	N72934	Hs.82614
467	245920	GYS2	glycogen synthase 2 (liver)	N52282	Hs.82614_
468		MGE1	GrpE-like protein cochaperone mRNA	AA887226	
469		GT	GT .	AI659294	
470		GBP1	Guanylate binding protein 1, interferon- inducible, 67kD	AA486849	Hs.62661
471		GBP1	Guanylate binding protein 1, interferon- inducible, 67kD	AA486850	Hs.62661
472		CLCN3	H. sapiens RNA for CLCN3	AA461332	Hs.174139
473		CLCN3	H. sapiens RNA for CLCN3	AA459750	Hs.174139
474		No Data	H.sapiens DAP-3 mRNA	N78611	
475		No Data	H.sapiens DAP-3 mRNA	W07332	Hs.74362
470	-	CLPP	H.sapiens mRNA for CLPP	W58658	
47		CLPP	H.sapiens mRNA for CLPP	W58337	Hs.74362
47		MCSP	H.sapiens mRNA for mitochondrial capsule selenoprotein	AA609976	Hs.111850
47		РМРСВ	H.sapiens mRNA for M-phase phosphoprotein, mpp11	N29844	Hs.184211
48		РМРСВ	H.sapiens mRNA for M-phase phosphoprotein, mpp11	N57262	Hs.184211
48		IDH3G	H.sapiens mRNA for NAD (H)-specific isocitrate dehydrogenase gamma subuni precursor	AA459380	Hs.75253

			H.sapiens mRNA for NAD (H)-specific		
		TDITAC	isocitrate dehydrogenase gamma subunit	AA459606	Hs.75253
482	810942	IDH3G		AA439000	115.75255
	500000	NUNTUE A 1	H.sapiens mRNA for NADH dehydrogenase	AA111999	Hs.74823
483	530282	NDUFA1	H.sapiens mRNA for NADH	AAIII	113.74025
404	520292	NDUFA1	dehydrogenase	AA083784	· Hs.74823
484	530282	NDOFAI	H.sapiens mRNA for orphan nuclear	12203701	110.7.1022
485	813815	NR1I3	hormone receptor	AA447727	Hs.83623
463	913613	141115	H.sapiens mRNA for orphan nuclear		
486	813815	NR1I3	hormone receptor	AA447889	Hs.83623
100	013015		H.sapiens mRNA for rho GDP-		
487	810959	ARHGDIA	dissociation Inhibitor 1	AA459400	Hs.159161
10/			H.sapiens mRNA for rho GDP-		
488	810959	ARHGDIA	dissociation Inhibitor 1	AA459625	Hs.159161
			H.sapiens mRNA for TIM17 preprotein		
489	506032	TIM17	translocase	AA708446	Hs.20716
490	814526	EST	H.sapiens seb4D mRNA	AA459363	
491	814526	EST	H.sapiens seb4D mRNA	AA459588	
492	814460	SURF5	H.sapiens SURF-5 mRNA	AA459247	Hs.78354
493	814460	SURF5	H.sapiens SURF-5 mRNA	AA459472	Hs.78354
			harakiri, BCL2-interacting protein		
494	1671299	HRK	(contains only BH3 domain)	AI083676	Hs.87247
		7.07	heat shock 10kD protein 1 (chaperonin	A A 07070C	TT- 202620
495	1417985	<u>EST</u>	heat shock 10kD protein 1 (chaperonin	AA878786	Hs.203620
496	1417985	EST	10)	AI791247	Hs.203620
497	1870305	HSPB2	heat shock 27kD protein 2	AI245337	Hs.78846
498	1354618	HSPF2	heat shock 40kD protein 2	AA830392	Hs.172847
499	755904	HSPA1A	HEAT SHOCK 70 KD PROTEIN 1	AA496544	Hs.8997
500	1604342	HSPA9B	heat shock 70kD protein 9B (mortalin-2)	AA987644	Hs.3069
100	100 15 12		HEAT SHOCK COGNATE 71 KD		
501	884719	HSPA10	PROTEIN	AA629567	Hs.180414
		**************************************	HEAT SHOCK PROTEIN HSP 90- ALPHA	H88540	Hs.180532
502	253009	HSPCA	HEAT SHOCK PROTEIN HSP 90-		HS.160332
503	253009	HSPCA	ALPHA	H88588	Hs.180532
504	824031	HSJ2	heat shock protein, DNAJ-like 2	AA490946	
505	824031	HSJ2	heat shock protein, DNAJ-like 2	AA491132	
506	1606894	HSF4	heat shock transcription factor 4	AA999776	Hs.75486
			hect (homologous to the E6-AP (UBE3A)		
			carboxyl terminus) domain and RCC1		
507	562101	HERC1	(CHC1)-like domain (RLD) 1	AA211496	Hs.255561
			hect (homologous to the E6-AP (UBE3A)		
			carboxyl terminus) domain and RCC1		
508		HERC1	(CHC1)-like domain (RLD) 1	AA211544	Hs.255561
509		HMOX1	heme oxygenase (decycling) 1	T71757	Hs.202833
510		HMOX1	heme oxygenase (decycling) 1	T71606	Hs.202833
511		HMOX2	Heme oxygenase (decycling) 2	AA626370	Hs.83853
512		HNF6A	hepatocyte nuclear factor 6, alpha	AA699732	Hs.73168
513		HK1	Hexokinase 1	AA485271	Hs.118625
514		HK1	Hexokinase 1	AA485272 AA811712	Hs.118625
515		C46157	HEXOKINASE D, LIVER ISOZYMES	H71868	+
516	211780	No Data	Hexosaminidase B (beta polypeptide)	11/1000	

517	211780	No Data	Hexosaminidase B (beta polypeptide)	H71081	
	767441	HEXB	hexosaminidase B (beta polypeptide)	AA417946	Hs.51043
518	767441	HEXB	hexosaminidase B (beta polypeptide)	AA418121	Hs.51043
520	970591	HMG1	High-mobility group (nonhistone chromosomal) protein 1	AA683085	Hs.189509
521	363103	HMG2	High-mobility group (nonhistone chromosomal) protein 2	AA019511	Hs.80684
522	363103	HMG2	High-mobility group (nonhistone chromosomal) protein 2	AA019203	Hs.80684
523	80924	HO3	histidyl-tRNA synthetase	T70104	Hs.77798
524	80924	HÖ3	histidyl-tRNA synthetase	T70037	Hs.77798
525	812246	HLCS	holocarboxylase synthetase (biotin- [proprionyl-Coenzyme A-carboxylase (ATP-hydrolysing)] ligase)	AA455043	Hs.79375
526	712577	HCCS	HOLOCYTOCHROME C-TYPE SYNTHETASE	AA281548	Hs.88859
527	712577	HCCS	HOLOCYTOCHROME C-TYPE SYNTHETASE	AA281549	Hs.88859
528	611075	HOXA1	homeo box A1	AA173231	
529	611075	HOXA1	homeo box A1	AA173290	
		APAF1	Homo sapiens apoptotic protease activating factor 1 (Apaf-1) mRNA, complete cds	N51014	Hs.77579
530	244146	APAF1	Homo sapiens apoptotic protease activating factor 1 (Apaf-1) mRNA, complete cds	N72045	Hs.77579
532	288796	BNIP2	Homo sapiens BCL2/adenovirus E1B 19kD-interacting protein 2 (BNIP2) mRNA, complete cds	N62514	Hs.155596
533	432620	NOD1	Homo sapiens caspase recruitment domain 4 (NOD1) mRNA	AA699441	
534	853506	23956 Mma	Homo sapiens clone 23956 mRNA, partial cds	AA663549	Hs.8128
535	288888	EST	Homo sapiens clone 640 unknown mRNA, complete sequence	N62617	Hs.44563
536	288888	EST	Homo sapiens clone 640 unknown mRNA, complete sequence	N78388	Hs.44563
537	271102	ccs	Homo sapiens copper chaperone for superoxide dismutase (CCS) mRNA, complete cds	N30404	Hs.5002
538	489823	COX17	Homo sapiens COX17 mRNA, complete cds	AA099855	Hs.16297
539	489823	COX17	Homo sapiens COX17 mRNA, complete cds	AA099320	Hs.16297
540	487348	DYMPLE	Homo sapiens dynamin-like protein mRNA, complete cds	AA045529	Hs.180628
541	487348	DYMPLE	Homo sapiens dynamin-like protein mRNA, complete cds	AA040651	Hs.180628
542	359982	BNIP3	Homo sapiens E1B 19K/Bcl-2-binding protein Nip3 mRNA, nuclear gene encoding mitochondrial protein, complete cds	AA063521	Hs.79428

			Homo sapiens E1B 19K/Bcl-2-binding protein Nip3 mRNA, nuclear gene		1
543	359982	BNIP3	encoding mitochondrial protein, complete	AA035669	Hs.79428
544	395436	HUEL	Homo sapiens embryonic lung protein (HUEL) mRNA, complete cds	AA757427	Hs.44053
545	1472184	HUEL	Homo sapiens embryonic lung protein (HUEL) mRNA, complete cds	AA873264	Hs.44053
546	129644	SSH3BP1	Homo sapiens eps8 binding protein e3B1 mRNA, complete cds	R16667	Hs.24752
547	129644	SSH3BP1	Homo sapiens eps8 binding protein e3B1 mRNA, complete cds Homo sapiens FRG1 mRNA, complete	R16666	Hs.24752
548	563574	FRG1	cds	AA113339	Hs.203772
549	563574	FRG1	Homo sapiens FRG1 mRNA, complete cds	AA112636	Hs.203772
	20052	TTD (02	Homo sapiens inner mitochondrial membrane translocase Tim23 (TIM23) mRNA, nuclear gene encoding	T69217	TT- 11000
550	83279	TIM23	mitochondrial protein, complete cds	T68317	Hs.11866
551	83279	TIM23	Homo sapiens inner mitochondrial membrane translocase Tim23 (TIM23) mRNA, nuclear gene encoding mitochondrial protein, complete cds	T68386	Hs.11866
		YZIA A 0.420	Homo sapiens KIAA0430 mRNA, complete cds	AA676805	Hs.77694
552	455123	KIAA0429	Homo sapiens mRNA for ATP binding	AA070803	HS.77094
553	300590	APACD	protein, complete cds	N80741	Hs.153884
554	300590	APACD	Homo sapiens mRNA for ATP binding protein, complete cds	W07537	Hs.153884
555	129146	COX7RP	Homo sapiens mRNA for COX7RP, complete cds	R10896	Hs.30888
556	129146	COX7RP	Homo sapiens mRNA for COX7RP, complete cds	R10947	Hs.30888
557	745314	HIRIP5	Homo sapiens mRNA for HIRA- interacting protein HIRIP5	AA625581	Hs.17368
558	810316	KIAA0631	Homo sapiens mRNA for KIAA0631 protein, partial cds	AA464206	Hs.75154
559	810316	KIAA0631	Homo sapiens mRNA for KIAA0631 protein, partial cds	AA464139	Hs.75154
560	490778	QP-C	Homo sapiens mRNA for low molecular mass ubiquinone-binding protein, complete cds	AA133191	Hs.3709
561	131653	RPMS12	Homo sapiens mRNA for mitochondrial ribosomal protein S12	R23752	1 Hs.9964
562	131653	RPMS12	Homo sapiens mRNA for mitochondrial ribosomal protein S12	R23806	Hs.9964
563	344825	LAS	Homo sapiens mRNA for putative lipoic acid synthetase, partial	W72965	Hs.53531
564		LAS	Homo sapiens mRNA for putative lipoic acid synthetase, partial	W76202	Hs.53531
565		CXX1	Homo sapiens mRNA for putatively prenylated protein	W72596	Hs.255532

ł	1		Homo sapiens mRNA for putatively	W76590	Hs.255532
566	345621	CXX1	prenylated protein	44 10230	110.633332
567	450574	SCA7	ataxia /	AA704255	Hs.108447
	123474	SCD	Homo_sapiens mRNA for stearoyl-CoA desaturase	R00707	Hs.119597
568		SCD	Homo sapiens mRNA for stearoyl-CoA desaturase	R00706	Hs.119597
569	123474 855890	cDNA	Homo sapiens mRNA; cDNA DKFZp586C0722 (from clone DKFZp586C0722)	AA630320	Hs.255914
570		NDUFV1	Homo sapiens NADH:ubiquinone dehydrogenase 51 kDa subunit (NDUFV1) mRNA, nuclear gene encoding mitochondrial protein, complete cds	AA427570	Hs.7744
571	770043	NDUFV1	Homo sapiens NADH:ubiquinone dehydrogenase 51 kDa subunit (NDUFV1) mRNA, nuclear gene encoding mitochondrial protein, complete cds	AA427652	Hs.7744
572	859228	IDH1	Homo sapiens NADP-dependent isocitrate dehydrogenase (IDH) mRNA, complete cds	AA666366	Hs.11223
573	124753	ABCD4	Homo sapiens peroxisomal membrane protein 69 (PMP69) mRNA, complete cds	R02189	Hs.94395
574 575	293104	РНҮН	Homo sapiens peroxisomal phytanoyl- CoA alpha-hydroxylase (PAHX) mRNA, complete cds	N91990	Hs.172887
	293104	рнүн	Homo sapiens peroxisomal phytanoyl- CoA alpha-hydroxylase (PAHX) mRNA, complete cds	N63845	Hs.172887
576		hTIM44	Homo sapiens putative mitochondrial inner membrane protein import receptor (hTIM44) mRNA, nuclear gene encoding mitochondrial protein, complete cds	AI003393	Hs.123178
577		RPA40	Homo sapiens RNA polymerase I subunit hRPA39 mRNA, complete cds	AA733038	Hs.5409
578 579		SLC4A4	Homo sapiens sodium bicarbonate cotransporter (HNBC1) mRNA, complete cds	AA452278	Hs.5462
		WFS1	Homo sapiens transmembrane protein (WFS1) mRNA, complete cds	R53910	Hs.26077
580		WFS1	Homo sapiens transmembrane protein (WFS1) mRNA, complete cds	R53911	Hs.26077
582		YWHAE	Human 14-3-3 epsilon mRNA, complete cds	N21624	Hs.79474
58:		YWHAE	Human 14-3-3 epsilon mRNA, complete cds	N28863	Hs.79474
58		AK2	Human adenylate kinase 2 (adk2) mRNA complete cds	H09730	Hs.17181
58		AK2	Human adenylate kinase 2 (adk2) mRNA complete cds	H10488	Hs.17181
58		AOE372	Human antioxidant enzyme AOE37-2 mRNA, complete cds	AA459663	Hs.83383

587	241481	CASP10	Human apoptotic cysteine protease Mch4 (Mch4) mRNA, complete cds	H80712	Hs.5353
588	241481	CASP10	Human apoptotic cysteine protease Mch4 (Mch4) mRNA, complete cds	H80711	Hs.5353
589	897774	APRT	Human APRT gene for adenine phosphoribosyltransferase	AA598510	Hs.28914
590	810552	REA	Human B-cell receptor associated protein (hBAP) mRNA, partial cds	AA464669	Hs.7771
591	810552	REA	Human B-cell receptor associated protein (hBAP) mRNA, partial cds	AA464567	Hs.7771
592	2326129	ввс3	Human Bel-2 binding component 3 (bbc3) mRNA, partial eds	AI688112	Hs.87246
593	795729	BAD	Human Bcl-2 binding component 6 (bbc6) mRNA, complete cds	AA460291	Hs.76366
594	795729	BAD	Human Bcl-2 binding component 6 (bbc6) mRNA, complete cds	AA461579	Hs.76366
595	814478	BCL2A1	Human Bcl-2 related (Bfl-1) mRNA, complete cds	AA459263	Hs.227817
596	814478	BCL2A1	Human Bcl-2 related (Bfl-1) mRNA, complete cds	AA459491	Hs.227817
597	212198	TP53BP2	Human Bcl2, p53 binding protein Bbp/53BP2 (BBP/53BP2) mRNA, complete cds	H69153	Hs.44585
			Human Bcl2, p53 binding protein Bbp/53BP2 (BBP/53BP2) mRNA, complete cds	H69077	Hs.44585
598	212198 432581	TP53BP2 Novel Gene	Human BRCA2 region, mRNA sequence CG012	AA699390	Hs.184938
600	433111	BAC clone CIT987SK-A- 735G6	Human Chromosome 16 BAC clone CIT987SK-A-735G6	AA680407	Hs.47278
601	782635	NDUFAB1	Human Chromosome 16 BAC clone CIT987SK-A-735G6	AA447569	Hs.5556
602	782635	NDUFAB1	Human Chromosome 16 BAC clone CIT987SK-A-735G6	AA448553	Hs.5556
603	669471	23589 mRNA	Human clone 23589 mRNA sequence	AA234889	Hs.11506
604	669471	23589 mRNA	Human clone 23589 mRNA sequence	AA253479	Hs.11506
605	771173	23732 mRNA	Human clone 23732 mRNA, partial cds	AA443497	Hs.81281
606	771173	23732 mRNA	Human clone 23732 mRNA, partial cds	AA429483	Hs.81281
607	745606	23759 mRNA	Human clone 23759 mRNA, partial cds	AA626336	Hs.118666
608	323500	CASP6	Human cysteine protease Mch2 isoform alpha (Mch2) mRNA, complete cds	W45688	Hs.3280
609	323500	CASP6	Human cysteine protease Mch2 isoform alpha (Mch2) mRNA, complete cds	W44316	Hs.3280
610	71420	CYP2C9	Human cytochrome P4502C9 (CYP2C9) mRNA, clone 65	T47787	
611	71420	CYP2C9	Human cytochrome P4502C9 (CYP2C9) mRNA, clone 65	T47788	
612	754490	SLC25A16	Human GT mitochondrial solute carrier protein homologue mRNA, complete cds	AA411554	Hs.180408
613	754490	SLC25A16	Human GT mitochondrial solute carrier protein homologue mRNA, complete cds	AA410572	Hs.180408

614	1734348	Go-alpha	Human guanine nucleotide-binding regulatory protein (Go-alpha) gene	AI188097	Hs.169647
-			Human homolog of D. melanogaster		
			flightless-I gene product mRNA, partial		
615	826204	FLII	cds	AA521453	Hs.83849
			Human HS1 binding protein HAX-1		
1			mRNA, nuclear gene encoding		
616	144777	HAX1	mitochondrial protein, complete cds	R76263	Hs.15318
			Human HS1 binding protein HAX-1		
			mRNA, nuclear gene encoding		
617	144777	HAX1	mitochondrial protein, complete cds	R76544	Hs.15318
618	261481	CUL3	Human Hs-cul-3 mRNA, partial cds	H98621	Hs.78946
619	261481	CUL3	Human Hs-cul-3 mRNA, partial cds	N25142	Hs.78946
			Human huntingtin interacting protein		
620	233581	HIP2	(HIP2) mRNA, complete cds	H78483	Hs.155485
			Human huntingtin interacting protein	-	
621	233581	HIP2	(HIP2) mRNA, complete cds	H78385	Hs.155485
			Human mitochondrial creatine kinase		
622	363086	CKMT1	(CKMT) gene, complete cds	AA019482	Hs.153998
			Human mitochondrial creatine kinase		- · · · · - ·
623	363086	CKMT1	(CKMT) gene, complete cds	AA019332	Hs.153998
			Human mitochondrial intermediate		
			peptidase precursor (MIPEP) mRNA,		
			mitochondrial gene encoding		
624	1390584	MIPEP	mitochondrial protein, complete cds	AA843592	Hs.68583
			Human mitochondrial RNA polymerase		
			mRNA, nuclear gene encoding		
625	134269	POLRMT	mitochondrial protein, complete cds	R31115	Hs.153880
			Human mitochondrial RNA polymerase		
			mRNA, nuclear gene encoding		
626	134269	POLRMT	mitochondrial protein, complete cds	R31174	Hs.153880
			Human mRNA for c-myc binding protein,		
627	781050	PFDN5	complete cds	AA446453	Hs.80686
			Human mRNA for c-myc binding protein,		
628	781050	PFDN5	complete cds	AA430010	Hs.80686
			Human mRNA for DNA polymerase		
629	451706	POLA	alpha-subunit	AA707650	Hs.74090
			Human mRNA for KIAA0017 gene,		
630	41541	EST	complete cds	R52789	Hs.253023
			Human mRNA for KIAA0017 gene,		
631	41541	EST	complete cds	R52788	Hs.253023
			Human mRNA for KIAA0070 gene,		
632	842818	KARS	partial cds	AA486374	Hs.3100
			Human mRNA for KIAA0070 gene,		
633	842818	KARS	partial cds	AA486220	Hs.3100
			Human mRNA for KIAA0123 gene,		
634	280496	KIAA0123	partial cds	N51632	Hs.75353
			Human mRNA for KIAA0185 gene,		
635	859627	KIAA0185	partial cds	AA666405	Hs.255573
			Human mRNA for KIAA0188 gene,		
636	783698	KIAA0188	partial cds	AA446822	Hs.81412
			Human mRNA for KIAA0188 gene,		
637	783698	KIAA0188	partial cds	AA446821	Hs.81412
ختت			X		

638	347031	KIAA0195	Human mRNA for KIAA0195 gene, complete cds	W79511	Hs.80540
038	347031		Human mRNA for KIAA0195 gene,		
639	347031	KIAA0195	complete cds	W79398	Hs.80540
			Human mRNA for KIAA0381 gene,		
640	753248	KIAA0381	partial cds	AA406231	Hs.100113
		•	Human mRNA for KIAA0381 gene,		
641	753248	KIAA0381	partial cds	AA406503	Hs.100113
			Human mRNA for mitochondrial 3-		
1			ketoacyl-CoA thiolase beta-subunit of	m/05/5	77 146010
642	108208	HADHB	trifunctional protein, complete cds	T69767	Hs.146812
			Human mRNA for mitochondrial 3-		
	100000	HADHB	ketoacyl-CoA thiolase beta-subunit of trifunctional protein, complete cds	T70752	Hs.146812
643	108208	палив	Human NADH:ubiquinone	170752	115.140012
			oxidoreductase MLRQ subunit mRNA,		
644	869538	NDUFA4	complete cds	AA680322	Hs.108661
			Human putative outer mitochondrial		
	•		membrane 34 kDa translocase hTOM34		
645	810452	TOM34	mRNA, complete cds	AA457118	Hs.76927
646	127197	CXORF6	Human Xq28 mRNA, complete cds	R08270	Hs.20136
647	127197	CXORF6	Human Xq28 mRNA, complete cds	R08271	Hs.20136
			Human zyxin related protein ZRP-1		1
648	811108	TRIP6	mRNA, complete cds	AA485677	Hs.119498
			Human zyxin related protein ZRP-1		·
649	811108	TRIP6	mRNA, complete cds	AA485807	Hs.119498
650	79828	HD	Huntingtin (Huntington disease)	T64094	Hs.79391
651	79828	HD	Huntingtin (Huntington disease)	T64015	Hs.79391
652	1752540	HIP1	huntingtin interacting protein 1	AI150389	
653	1566230	HYAL3	hyaluronoglucosaminidase 3	AI140794	Hs.129910
			HYDROXYACYLGLUTATHIONE	17100610	
654	2095653	GLO4	HYDROLASE	AI420642	
CEE	141966	HSD3B1	hydroxy-delta-5-steroid dehydrogenase, 3 beta- and steroid delta-isomerase 1	R68803	Hs.38586
655	141900	1100201	hydroxy-delta-5-steroid dehydrogenase, 3	100005	113.50500
656	141966	HSD3B1	beta- and steroid delta-isomerase 1	R68906	Hs.38586
657	126243	HMBS	Hydroxymethylbilane synthase	R06321	Hs.82609
658	126243	HMBS	Hydroxymethylbilane synthase	R06263	Hs.82609
659	321163	HSU79253	hypothetical protein	W53000	Hs.56155
660	321163	HSU79253	hypothetical protein	W52999	Hs.56155
000		*	Hypoxanthine phosphoribosyltransferase		
661	280507	HPRT1	1 (Lesch-Nyhan syndrome)	N47312	Hs.82314
			Hypoxanthine phosphoribosyltransferase		
662	280507	HPRT1	1 (Lesch-Nyhan syndrome)	N47311	Hs.82314
663	2216141	IFM1	IFM1	AI654459	
664	767034	ILVBL	ilvB (bacterial acetolactate synthase)-like	AA424288	Hs.78880
665	767034	ILVBL	ilvB (bacterial acetolactate synthase)-like	AA451741	Hs.78880
			IMP (inosine monophosphate)		
666	1606837	IMPDH2	dehydrogenase 2	AA996028	Hs.75432
			INITIATION FACTOR IF-2,		1
667	50754	MTIF2	MITOCHONDRIAL PRECURSOR	H18070	Hs.149894
			interferon, alpha-inducible protein (clone		
668	111981	G1P3	IFI-6-16)	T84633	Hs.179972

1	1		interferon, alpha-inducible protein (clone	T91807	Hs.179972
569	111981	G1P3	IFI-6-16)		Hs.126256
570	491763	IL1B	Interleukin 1, beta		Hs.126256
571	491763	IL1B	Interleukin 1, beta		HS.120230
572	2119594	IDH1	ISOCITRATE DEHYDROGENASE	AI399657	
673	869375	IDH2	Isocitrate dehydrogenase 2 (NADP+), mitochondrial	AA679907	Hs.252546
5/3	003070		isocitrate dehydrogenase 3 (NAD+) alpha		1
674	2168168	ETFB	(IDH3A),	AI565774	
675	755474	IARS	isoleucine-tRNA synthetase	AA410636	
676	755474	IARS	isoleucine-tRNA synthetase	AA419192	
577	810325	IVD	isovaleryl Coenzyme A dehydrogenase	AA464149	Hs.77510
678	810325	IVD	isovaleryl Coenzyme A dehydrogenase	AA464216	Hs.77510
679	2148505	JC4913	JC4913	AI469831	
	809707	JUNB	jun B proto-oncogene	AA454711	Hs.198951
680 681	809707	JUNB	jun B proto-oncogene	AA456366	Hs.198951
682	1474284	KAI1	kangai 1 (suppression of tumorigenicity 6, prostate; CD82 antigen (R2 leukocyte antigen, antigen detected by monoclonal and antibody IA4))	AA922309	Hs.25409
683	845441	KIAA0016	KIAA0016 gene product	AA644550	Hs.75187
684	2241563	KIHUA3	KIHUA3	AI632351	
685	21738	HADHSC	L-3-hydroxyacyl-Coenzyme A dehydrogenase, short chain	T65482	Hs.7153
003	21750		L-3-hydroxyacyl-Coenzyme A		
686	21738	HADHSC	dehydrogenase, short chain	T65407	Hs.7153
687	884644	LAMR1	Laminin receptor (2H5 epitope)	AA629897	Hs.181357
688	704360	LCFA HUMAN	LCFA HUMAN	AA279565	
689	1256792	MAD4	Mad4 homolog	AA875977	Hs.102402
690	1908746	MDH1	MALATE DEHYDROGENASE	AI302237	
691	1636908	MDH2	malate dehydrogenase 2, NAD (mitochondrial)	AI000271	Hs.111076
	2010949	ME2	malic enzyme 2, mitochondrial	AI361039	Hs.75342
692		ME3	malic enzyme, NADP+-dependent,	AA779401	Hs.2838
693	896921		Mannose-6-phosphate receptor (cation dependent)	AA465223	Hs.75709
694	814211	M6PR	Mannose-6-phosphate receptor (cation	1-1.00225	1
695	814211	M6PR	dependent)	AA465578	Hs.75709
696	589115	MMP1	matrix metalloproteinase 1 (interstitial collagenase)	AA143201	
697		MMP1	matrix metalloproteinase 1 (interstitial collagenase)	AA143331	
698		MMP11	matrix metalloproteinase 11 (stromelysin 3)	AA954935	Hs.155324
699		MMP12	matrix metalloproteinase 12 (macrophag elastase)	R92994	Hs.1695
70		MMP12	matrix metalloproteinase 12 (macrophag elastase)	e R93037	Hs.1695
		MMP17	matrix metalloproteinase 17 (membrane inserted)	н97792	Hs.159581
70		MMP15	Matrix metalloproteinase 2	AA443300	
70			Matrix metalloproteinase 2	AA443433	
70	3 784589	MMP15	Iviativ ilictatioproteinaso 2	1	

704	470393	MMP7	matrix metalloproteinase 7 (matrilysin, uterine)	AA031513	Hs.2256
705	470393	MMP7	matrix metalloproteinase 7 (matrilysin, uterine)	AA031514	Hs.2256
706	2213770	CCHU	Matrix metalloproteinase 7 (matrilysin, uterine)	AI582304	
707	22040	ммр9	matrix metalloproteinase 9 (gelatinase B, 92kD gelatinase, 92kD type IV collagenase)	T72581	
708	22040	ммр9	matrix metalloproteinase 9 (gelatinase B, 92kD gelatinase, 92kD type IV collagenase)	T64837	
709	773266	ммр9	matrix metalloproteinase 9 (gelatinase B, 92kD gelatinase, 92kD type IV collagenase)	AA425227	Hs.151738
710	773266	ММР9	matrix metalloproteinase 9 (gelatinase B, 92kD gelatinase, 92kD type IV collagenase)	AA425503	Hs.151738
711	2091860	MMPL1	matrix metalloproteinase-like 1	AI382081	Hs.198265
712	240766	Data unavailable	METALLOPROTEINASE INHIBITOR 1 PRECURSOR	H80214	
713	240766	Data unavailable	METALLOPROTEINASE INHIBITOR 1 PRECURSOR	H80215	
714	2165301	DEHUMT	methylene tetrahydrofolate dehydrogenase (NAD+ dependent), methenyltetrahydrofolate cyclohydrolase	AI497794	
715	1473506	MMSA_HUMAN	METHYLMALONATE- SEMIALDEHYDE DEHYDROGENASE	AA915881	
716	2091562	S40622	METHYLMALONYL-COA MUTASE PRECURSOR	AI377286	
717	278570	MITF	Microphthalmia-associated transcription factor	N66177	Hs.166017
718	278570	MITF	Microphthalmia-associated transcription factor	N99168	Hs.166017
719	768443	MGST1	microsomal glutathione S-transferase 1	AA495936	Hs.790
720	768443 629896	MGST1 MAP1B	microsomal glutathione S-transferase 1 MICROTUBULE-ASSOCIATED PROTEIN 1B	AA495935 AA219045	Hs.790 Hs.103042
721	1084029	A56650	MITOCHONDRIAL 2- OXOGLUTARATE/MALATE CARRIER PROTEIN	AA594052	
723		RPML3	MITOCHONDRIAL 60S RIBOSOMAL PROTEIN L3	H06113	Hs.79086
724		RPML3	MITOCHONDRIAL 60S RIBOSOMAL PROTEIN L3	H05820	Hs.79086
725	433553	CA5	Mitochondrial carbonic anhydrase	AA699469	Hs.177446
70.6	1175520	A39018	MITOCHONDRIAL CARNITINE PALMITOYLTRANSFERASE II PRECURSOR	AA641442	
726		TSFM	MITOCHONDRIAL ELONGATION FACTOR TS PRECURSOR	W47015	Hs.3273
728		TSFM	MITOCHONDRIAL ELONGATION FACTOR TS PRECURSOR	W47014	Hs.3273

١			MITOCHONDRIAL LON PROTEASE HOMOLOG	AI377406	
729	2066585	LON-PEN	MITOCHONDRIAL MATRIX PROTEIN P1 PRECURSOR	R26234	
730	133099	MMP1 Precursor MMP1 Precursor	MITOCHONDRIAL MATRIX PROTEIN P1 PRECURSOR	R26235	
731	448491	mrp2	MITOCHONDRIAL RIBOSOMAL PROTEIN S14	AA777627	
733	1587031	HSPA9	MITOCHONDRIAL STRESS-70 PROTEIN	AA948223	
734	814346	MTIF2	mitochondrial translational initiation factor 2	AA458825	Hs.149894
735	814346	MTIF2	mitochondrial translational initiation factor 2	AA459016	Hs.149894
736	1916771	MTRF1	mitochondrial translational release factor	AI347695	Hs.80683
737	1880757	MAP3K5	mitogen-activated protein kinase kinase kinase 5	AI268273	Hs.151988
738	2244621	MRF1	MRF1	AI656905	
739	645006	MRP4	MRP4	AA197284	
740	645006	MRP4	MRP4	AA205815	
741	1933324	MPZ	myelin protein zero (Charcot-Marie- Tooth neuropathy 1B)	AI351956	Hs.93883
	302190	MLL	myeloid/lymphoid or mixed-lineage leukemia (trithorax (Drosophila) homolog)	W16724	Hs.199160
742	302190	WILD	myeloid/lymphoid or mixed-lineage		
743	302190	MLL	leukemia (trithorax (Drosophila) homolog)	N77807	Hs.199160
744		MYL6	MYOSIN LIGHT CHAIN ALKALI, NON-MUSCLE ISOFORM	AA488346	Hs.77385
745		MYL6	MYOSIN LIGHT CHAIN ALKALI, NON-MUSCLE ISOFORM	AA488477	Hs.77385
746		MTHFD2	NAD-dependent methylene tetrahydrofolate dehydrogenase cyclohydrolase	AA480995	Hs.154672
		MTHFD2	NAD-dependent methylene tetrahydrofolate dehydrogenase cyclohydrolase	AA480994	Hs.154672
747		NDUFA2	NADH dehydrogenase (ubiquinone) 1 alpha subcomplex, 2 (8kD, B8)	AA425211	Hs.163867
749		NDUFA2	NADH dehydrogenase (ubiquinone) 1 alpha subcomplex, 2 (8kD, B8)	AA425534	Hs.163867
750		NDUFA2	NADH dehydrogenase (ubiquinone) 1 alpha subcomplex, 2 (8kD, B8)	AI017426	Hs.163867
75		NDUFA5	NADH dehydrogenase (ubiquinone) 1 alpha subcomplex, 5 (13kD, B13)	AA608515	Hs.83916
75		. NDUFA9	NADH dehydrogenase (ubiquinone) 1 alpha subcomplex, 9 (39kD)	AA598884	Hs.75227
75		NDUFB4	NADH dehydrogenase (ubiquinone) 1 beta subcomplex, 4 (15kD, B15)	AA704675	
75		NDUFS5	NADH dehydrogenase (ubiquinone) 1 beta subcomplex, 4 (15kD, B15)	AA214053	Hs.80595

755	562409	NDUFS5	NADH dehydrogenase (ubiquinone) 1 beta subcomplex, 4 (15kD, B15)	AA214154	Hs.80595
756	307933	NDUFB5	NADH dehydrogenase (ubiquinone) 1	•	
130	301933	Ирогра	beta subcomplex, 5 (16kD, SGDH)	N93053	Hs.19236
757	307933	NDUFB5	NADH dehydrogenase (ubiquinone) 1 beta subcomplex, 5 (16kD, SGDH)	W21390	Hs.19236
758	470861	NDUFB6	NADH dehydrogenase (ubiquinone) 1 beta subcomplex, 6 (17kD, B17)	AA034268	Hs.109646
759	470861	NDUFB6	NADH dehydrogenase (ubiquinone) 1 beta subcomplex, 6 (17kD, B17)	AA032077	Hs.109646
760	771089	NDUFB7	NADH dehydrogenase (ubiquinone) 1 beta subcomplex, 7 (18kD, B18)	AA428058	Hs.661
761	771089	NDUFB7	NADH dehydrogenase (ubiquinone) 1 beta subcomplex, 7 (18kD, B18)	AA429046	Hs.661
1			NADH dehydrogenase (ubiquinone) 1,	211123040	113.001
762	796513	NDUFC1	subcomplex unknown, 1 (6kD, KFYI)	AA460251	Hs.84549
763	796513	NDUFC1	NADH dehydrogenase (ubiquinone) 1, subcomplex unknown, 1 (6kD, KFYI)	AA463815	Hs.84549
764	753457	NDUFS1	NADH dehydrogenase (ubiquinone) Fe-S protein 1 (75kD) (NADH-coenzyme Q reductase)	AA406535	Hs.8248
70.	700.07	N. DOTE	NADH dehydrogenase (ubiquinone) Fe-S protein 1 (75kD) (NADH-coenzyme Q	AA400333	118.0248
765	753457	NDUFS1	reductase)	AA406536	Hs.8248
766	743811	NDUFS3	NADH dehydrogenase (ubiquinone) Fe-S protein 3 (30kD) (NADH-coenzyme Q reductase)	AA634381	Hs.5273
767	377152	NDUFS4	NADH dehydrogenase (ubiquinone) Fe-S protein 4 (18kD) (NADH-coenzyme Q reductase)	AA055101	Hs.10758
768	377152	NDUFS4	NADH dehydrogenase (ubiquinone) Fe-S protein 4 (18kD) (NADH-coenzyme Q reductase)	AA055102	Hs.10758
769	502141	NDUFS8	NADH dehydrogenase (ubiquinone) Fe-S protein 8 (23kD) (NADH-coenzyme Q reductase)	AA127014	Hs.90443
770	502141	NDUFS8	NADH dehydrogenase (ubiquinone) Fe-S protein 8 (23kD) (NADH-coenzyme Q reductase)	AA128218	Hs.90443
771	1486260	NDUFV2	NADH dehydrogenase (ubiquinone) flavoprotein 2 (24kD)		
772	1676983	RDHUB5	NADH-CYTOCHROME B5 REDUCTASE	AA922326 AI076798	Hs.51299
			natriuretic peptide receptor B/guanylate cyclase B (atrionatriuretic peptide		
773	1632011	NPR2	receptor B)	AA994689	Hs.78518
774	994950	NDUFA10	NDUFA10	AA555087	
775	1588691	NDUFA2	NDUFA2	AA975360	
776	929358	NDUFA3	NDUFA3	AA492039	
777	2118546	NDUFA6	NDUFA6	AI524304	
778	1270489	NDUFA7	NDUFA7	AA748476	
779	171756	NDUFA8	NDUFA8	H18333	

818		PPIF	peptidylprolyl isomerase F (cyclophilin	AA442184	Hs.173125
817	774726	PPIF	F)	AA442081	Hs.173125
816		PPIF	PRECURSOR peptidylprolyl isomerase F (cyclophilin	H05115	Hs.173125
			PEPTIDYL-PROLYL CIS-TRANS ISOMERASE, MITOCHONDRIAL		Y 15010-
815	43884	PPIF	ISOMERASE, MITOCHONDRIAL PRECURSOR	H05580	Hs.173125
014	102773	2020	PEPTIDYL-PROLYL CIS-TRANS		
813 814		DDX5	P68 PROTEIN	H27646	Hs.76053
812		UQCRC1 DDX5	PROTEIN I PRECURSOR P68 PROTEIN	AI005342 H27564	Hs.76053
			P31930 UBIQUINOL-CYTOCHROME- C REDUCTASE COMPLEX CORE		
811		OGDH	oxoglutarate dehydrogenase (lipoamide)	AA856769	Hs.168669
810	233479	No Data	Oxoglutarate dehydrogenase (lipoamide)	H80138	
809	233479	No Data	Oxoglutarate dehydrogenase (lipoamide)	H78910	
808	1704170	OXA1HS_	oxoglutarate dehydrogenase (lipoamide)	AI096611	
807	1637751	OXAIL	oxidase (cytochrome c) assembly 1-like	AI001180	Hs.151134
806	796646	ODC1	Ornithine decarboxylase 1	AA461467	Hs.75212
805	796646	ODC1	Ornithine decarboxylase 1	AA460115	Hs.75212
804	783696	OAT	atrophy)	AA446820	Hs.75485
803	783696	OAT	atrophy) ornithine aminotransferase (gyrate	AA446819	Hs.75485
802	30/414	TALAD	ornithine aminotransferase (gyrate		
801	1870662 567414	NFYB	nuclear transcription factor Y, beta	AA130846	
800	207808	NFATC3 A54868	nuclear factor of activated T-cells, cytoplasmic 3 nuclear respiratory factor 1 (NRF1)	H59047 AI245773	Hs.172674
799	207808	NFATC3	nuclear factor of activated T-cells, cytoplasmic 3	H59048	Hs.172674
798	755750	NME2	non-metastatic cells 2, protein (NM23B) expressed in	AA496512	Hs.255790
797	755750	NME2	expressed in	AA496628	Hs.255790
			non-metastatic cells 2, protein (NM23B)	A A 406609	Ша 255700
796	1468820	NOS3	nitric oxide synthase 3 (endothelial cell)	AA884967	Hs.166373
795	150466	EST	Nitric oxide synthase 3 (endothelial cell)	H01788	Hs.237163
794	150466	EST	Nitric oxide synthase 3 (endothelial cell)	H01039	Hs.237163
793	1160732	NOS2A	Nitric oxide synthase 2A (inducible, hepatocytes)	AA877840	Hs.193788
792	51826	NNT	nicotinamide nucleotide transhydrogenase	H24126	Hs.18136
791	51826	NNT	nicotinamide nucleotide transhydrogenase	H22944	Hs.18136
790	1585499	NDUFS5	NDUFS5	AA974058	
789	1019401	NDUFS2	NDUFS2	AA551149	
788	2277817	NDUFC2	NDUFC2	AI653801	
787	984496	NDUFB9	NDUFB9	AA526075	
785 786	2266937	NDUFB8	NDUFB8	AI608733	
784	2103675	NDUFB3	NDUFB3	AI401200	
783	2170527 1881693	NDUFB2	NDUFB2	AI290799	
782	998616	NDUFB1 NDUFB10	NDUFB10	AI571333	
781	1473135	NDUFAB1	NDUFAB1	AA535762	
		2 TOTAL	NDUFAB1	AA873566	

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819	133273	PMP22	peripheral myelin protein 22	R26960	Hs.103724
820	133273	PMP22	peripheral myelin protein 22	R26732	Hs.103724
			Peroxisomal membrane protein 3 (35kD,		
821	788518	PXMP3	Zellweger syndrome)	AA452566	
			Peroxisomal membrane protein 3 (35kD,		
822	788518	PXMP3	Zellweger syndrome)	AA452747	
823	1476157	PXR1	peroxisome receptor 1	AA873073	Hs.158084
824	455263	PET112L	PET112 (yeast homolog)-like	AA677572	Hs.11127
825	951683	PHC	PHC	AA629980	
826	843109	FARS1	phenylalanine-tRNA synthetase	AA488691	Hs.57969
827	843109	FARS1	phenylalanine-tRNA synthetase	AA486503	Hs.57969
828	842784	PHC	phosphate carrier, mitochondrial	AA486305	Hs.78713
829	842784	PHC	phosphate carrier, mitochondrial	AA486200	Hs.78713
830	346009	PFKL	Phosphofructokinase (liver type)	W72140	Hs.155455
831	346009	PFKL	Phosphofructokinase (liver type)	W77881	Hs.155455
832	489626	PFKM	Phosphofructokinase, muscle	AA099169	Hs.75160
833	489626	PFKM	Phosphofructokinase, muscle	AA101919	Hs.75160
834	183194		phospholipase A2, group IIA (platelets, synovial fluid)	H44953	
835	183194	_	phospholipase A2, group IIA (platelets, synovial fluid)	H45000	
836	77915	PLA2G2A	phospholipase A2, group IIA (platelets, synovial fluid)	T61323	Hs.76422
			phospholipase A2, group IIA (platelets,		
837	77915	PLA2G2A	synovial fluid)	T61271	Hs.76422
838	2367279	PN0673	PN0673	AI741984	
839	780947	POLD1	Polymerase (DNA directed), delta 1, catalytic subunit (125kD)	AA429661	Hs.99890
840	780947	POLD1	Polymerase (DNA directed), delta 1, catalytic subunit (125kD)	AA446151	Hs.99890
841	626206	POLG	polymerase (DNA directed), gamma	AA188761	Hs.80961
842	626206	POLG	polymerase (DNA directed), gamma	AA188629	Hs.80961
843	827129	POLRMT	polymerase (RNA) mitochondrial (DNA directed)	AA521239	Hs.153880
844	2144562	PRAX-1	PRAX-1	AI452994	
845	826211	PDCD2	Programmed cell death 2	AA521466	Hs.41639
846	814337	PCCB	propionyl Coenzyme A carboxylase, beta polypeptide	AA459122	Hs.63788
			propionyl Coenzyme A carboxylase, beta		
847	814337	PCCB	polypeptide	AA459341	Hs.63788
848	950710	PCCA	Propionyl-coA carboxylase alpha chain	AA608575	Hs.80741
849	824568	KLK3	Prostate specific antigen	AA490981	Hs.171995
850	843134	PBP	Prostatic binding protein	AA486514	Hs.80423
851	843134	PBP	Prostatic binding protein	AA485909	Hs.80423
852	261971	PRSM1	protease, metallo, 1, 33kD	N27227	Hs.183138
853	261971	PRSM1	protease, metallo, 1, 33kD	H98666	Hs.183138
854	375728	PRSM1	protease, metallo, 1, 33kD	AA033807	Hs.183138
855	51041	Est	protoporphyrinogen oxidase	H18633	Hs.123641
856	51041	Est	protoporphyrinogen oxidase	H18716	Hs.123641
857	247160	PPOX	protoporphyrinogen oxidase	N57891	Hs.100016
858	504452	PPOX	protoporphyrinogen oxidase	AA151249	Hs.100016
859	504452	PPOX	protoporphyrinogen oxidase	AA151248	Hs.100016

860	501939	HHCMA56	putative oxidoreductase	AA128041	Hs.519
861	501939	ННСМА56	putative oxidoreductase	AA128086	Hs.519
862	108378	PC	pyruvate carboxylase	T77729	Hs.89890
863	108378	PC	pyruvate carboxylase	T77728	Hs.89890
			Pyruvate dehydrogenase (lipoamide)		
864	80374	PDHA1	alpha 1	T65833	Hs.1023
			Pyruvate dehydrogenase (lipoamide)		
865	80374	PDHA1	alpha 1	T65758	Hs.1023
866	826077	PDHB	pyruvate dehydrogenase (lipoamide) beta	AA521401	Hs.979
867	279665	PDX1	Pyruvate dehydrogenase complex, lipoyl- containing component X; E3-binding protein	N48320	Hs.74642
868	279665	PDX1	Pyruvate dehydrogenase complex, lipoyl- containing component X; E3-binding protein	N49046	Hs.74642
808	279003	10/11	PYRUVATE DEHYDROGENASE E1	1149040	113.74042
869	1542260	PDHA2	COMPONENT, ALPHA	AA927400	
870	1645668	PDK1	pyruvate dehydrogenase kinase, isoenzyme 1	AI026814	Hs.159477
871	66686	No Data	QM PROTEIN	T67270	
872	66686	No Data	QM PROTEIN	T67271	
873	590727	RENT1	regulator of nonsense transcripts 1	AA156342	Hs.12719
874	590727	RENT1	regulator of nonsense transcripts 1	AA156376	Hs.12719
875	624627	RRM2	Ribonucleotide reductase M2 polypeptide	AA187351	Hs.75319
876	624627	RRM2	Ribonucleotide reductase M2 polypeptide	AA188430	Hs.75319
877	788334	RPL23L	ribosomal protein L23-like	AA453015	Hs.3254
878	730124	RPL7	Ribosomal protein L7	AA412470	Hs.153
879	730124	RPL7	Ribosomal protein L7	AA412344	Hs.153
880	809578	RPS5	Ribosomal protein S5	AA456616	Hs.76194
881	809578	RPS5	Ribosomal protein S5	AA455795	Hs.76194
882	214133	FSRG1	RING3 PROTEIN	H72520	Hs.75243
883	214133	FSRG1	RING3 PROTEIN	H72918	Hs.75243
			SCO (cytochrome oxidase deficient,		
884	110744	SCO2	yeast) homolog 2	T90560	Hs.180903
885	110744	SCO2	SCO (cytochrome oxidase deficient, yeast) homolog 2	T83097	Hs.180903
886	646657	SCO1	SCO1 (yeast homolog) cytochrome oxidase deficient 1	AA205413	Hs.14511
887	646657	SCO1	SCO1 (yeast homolog) cytochrome oxidase deficient 1	AA205579	Hs.14511
888	1074487	SDHD	SDHD	AA579646	
889	951117	SHMT2	serine hydroxymethyltransferase 2 (mitochondrial)	AA620477	
890	2138479	XNHUSP	SERINEPYRUVATE AMINOTRANSFERASE	AI521720	
891	125183	SSBP	single-stranded DNA-binding protein	R05693	Hs.923
892	125183	SSBP	single-stranded DNA-binding protein	R05694	Hs.923
893	813678	SLC1A3	solute carrier family 1 (glial high affinity glutamate transporter), member 3	AA453823	Hs.75379
894	813678	SLC1A3	solute carrier family 1 (glial high affinity glutamate transporter), member 3	AA453742	Hs.75379
895	586990	SLC11A2	solute carrier family 11 (proton-coupled divalent metal ion transporters), member	AA133656	Hs.57435

			2		
896	586990	SLC11A2	solute carrier family 11 (proton-coupled divalent metal ion transporters), member	AA133655	Hs.57435
897	2029995	SLC12A7	solute carrier family 12 (potassium/chloride transporters), member 7	AI492956	Hs.172613
898	190732	SLC2A5	Solute carrier family 2 (facilitated glucose transporter), member 5	H38650	Hs.33084
899	190732	SLC2A5	Solute carrier family 2 (facilitated glucose transporter), member 5	Н38733	Hs.33084
900	758304	SLC25A14	solute carrier family 25 (mitochondrial carrier, brain), member 14	AA404241	Hs.194686
901	758304	SLC25A14	solute carrier family 25 (mitochondrial carrier, brain), member 14	AA401224	Hs.194686
902	897107	SLC25A1	solute carrier family 25 (mitochondrial carrier; citrate transporter), member 1	AA676877	Hs.111024
903	878413	SLC25A11	solute carrier family 25 (mitochondrial carrier; oxoglutarate carrier), member 11	AA670357	Hs.184877
904	433350	SORD	Sorbitol dehydrogenase	AA700604	Hs.878
905	1256737	SPG7	spastic paraplegia 7, paraplegin (pure and complicated autosomal recessive)	AA876165	Hs.78497
906	682528	SCA1	spinocerebellar ataxia 1 (olivopontocerebellar ataxia 1, autosomal dominant, ataxin 1)	AA256507	Hs.74520
907	682528	SCA1	spinocerebellar ataxia 1 (olivopontocerebellar ataxia 1, autosomal dominant, ataxin 1)	AA256508	Hs.74520
908	128875	SCA2	Spinocerebellar ataxia 2 (olivopontocerebellar ataxia 2, autosomal dominant)	R10604	Hs.76253
909	128875	SCA2	Spinocerebellar ataxia 2 (olivopontocerebellar ataxia 2, autosomal dominant)	R10603	Hs.76253
910	447167	SFRS8	splicing factor, arginine/serine-rich 8 (suppressor-of-white-apricot, Drosophila homolog)	AA702973	Hs.84229
911	1934125	B34181	steroid 11-beta-hydroxylase	AI334352	
912	859858	STAR	steroidogenic acute regulatory protein	AA679454	Hs.3132
913	855395	SCP2	sterol carrier protein 2	AA664009	Hs.75760
914	824699	SUCA_HUMAN	SUCA_HUMAN	AA482206	
915	80915	SDHA	Succinate dehydrogenase 2, flavoprotein (Fp) subunit	T70043	
916	80915	SDHA	Succinate dehydrogenase 2, flavoprotein (Fp) subunit	T70109	
917	797016	SDHB	succinate dehydrogenase complex, subunit B, iron sulfur (Ip)	AA463565	Hs.64
918	797016	SDHB	succinate dehydrogenase complex, subunit B, iron sulfur (Ip)	AA463510	Hs.64
919	366132	SDHC	succinate dehydrogenase complex, subunit C, integral membrane protein, 15kD	AA062805	Hs.3577

920	504145	LSC2	SUCCINATE-COA LIGASE	AA132000	
921	504145	LSC2	SUCCINATE-COA LIGASE	AA131817	
922	845630	SUCLA2	succinate-CoA ligase, ADP-forming, beta subunit	AA644577	Hs.182217
923	1571111	SCOT HUMAN	Succinyl CoA:3-oxoacid CoA transferase	AA936982	
924	950489	SOD1	Superoxide dismutase 1 (Cu/Zn)	AA599127	Hs.75428
925	1572467	SOD2	Superoxide dismutase 2	AA934677	
926	346860	SOD2	superoxide dismutase 2, mitochondrial	W78148	Hs.177781
926	346860	SOD2	superoxide dismutase 2, mitochondrial	W79913	Hs.177781
	840708	SOD2	Superoxide dismutase 2, mitochondrial	AA488084	Hs.177781
928	840708	SOD2	Superoxide dismutase 2, mitochondrial	AA487750	Hs.177781
929	1343732	SOD3	Superoxide dismutase 3, extracellular	AA725564	Hs.2420
930	433474	SURF1	Surfeit 1	AA699560	Hs.3196
931	2114302	surfeit 1	surfeit 1	AI417847	Hs.3196
932	782797	SMN1	survival of motor neuron 1, telomeric	AA448194	Hs.77306
933	704299	TAZ	tafazzin (cardiomyopathy, dilated 3A (X- linked); endocardial fibroelastosis 2; Barth syndrome)	AA279440	Hs.79021
935	789376	TXNRD1	thioredoxin reductase 1	AA453335	Hs.13046
936	789376	TXNRD1	thioredoxin reductase 1	AA464849	Hs.13046
937	796000	MPST	thiosulfate sulfurtransferase (rhodanese)	AA461065	Hs.74097
938	796000	MPST	thiosulfate sulfurtransferase (rhodanese)	AA460495	Hs.74097
939	856167	TARS	threonyl-tRNA synthetase	AA630628	Hs.84131
940	205185	THBD	Thrombomodulin	H59861	Hs.2030
941	205185	THBD	Thrombomodulin	H60674	Hs.2030
		TIAL 1	TIA1 cytotoxic granule-associated RNA- binding protein-like 1	N59426	Hs.182741
942	246300	TIAL1 TIMP2	tissue inhibitor of metalloproteinase 2	AA486280	Hs.246948
943	842846 489519	TIMP3	tissue inhibitor of metalloproteinase 3 (Sorsby fundus dystrophy, pseudoinflammatory)	AA099153	Hs.245188
945		TIMP3	tissue inhibitor of metalloproteinase 3 (Sorsby fundus dystrophy, pseudoinflammatory)	AA099251	Hs.245188
946	754106	TIMP3	tissue inhibitor of metalloproteinase 3 (Sorsby fundus dystrophy, pseudoinflammatory)	AA479202	Hs.245188
947	754106	TIMP3	tissue inhibitor of metalloproteinase 3 (Sorsby fundus dystrophy, pseudoinflammatory)	AA478662	Hs.245188
948	433481	TCF2	transcription factor 2, hepatic; LF-B3; variant hepatic nuclear factor	AA699573	Hs.169853
949		TCF6L1	Transcription factor 6-like 1 (mitochondrial transcription factor 1-like	AA449118	Hs.75133
950	785845	TCF6L1	Transcription factor 6-like 1 (mitochondrial transcription factor 1-like	AA449551	Hs.75133
951	841334	STIP1	TRANSFORMATION-SENSITIVE PROTEIN IEF SSP 3521	AA487635	Hs.75612
952		STIP1	TRANSFORMATION-SENSITIVE PROTEIN IEF SSP 3521	AA487427	Hs.75612
95:	3 2114841	U66035	translocase of inner mitochondrial membrane 8	AI432207	Hs.102824
95	4 970880	TPM4	tropomyosin 4	AA774983	1113.102024

955	341328	TPM1	Tropomyosin alpha chain (skeletal muscle)	W58092	Hs.77899
956	341328	TPM1	Tropomyosin alpha chain (skeletal muscle)	W58009	Hs.77899
957	32621	WARS2	tryptophanyl tRNA synthetase 2 (mitochondrial)	R43272	Hs.227274
958	32621	WARS2	tryptophanyl tRNA synthetase 2 (mitochondrial)	R18903	Hs.227274
959	684582	WARS2	tryptophanyl tRNA synthetase 2 (mitochondrial)	AA251354	Hs.227274
960	684582	WARS2	tryptophanyl tRNA synthetase 2 (mitochondrial)	AA251468	Hs.227274
961	594079	TID1	tumorous imaginal discs (Drosophila) homolog	AA169872	Hs.6216
962	594079	TID1	tumorous imaginal discs (Drosophila) homolog	AA169567	Hs.6216
963	1569989	YWHAB	tyrosine 3-monooxygenase/tryptophan 5- monooxygenase activation protein, beta polypeptide	AA962407	Hs.182238
964	1591788	YWHAZ	tyrosine 3-monooxygenase/tryptophan 5- monooxygenase activation protein, zeta polypeptide	AA976477	Hs.75103
965	25499	UQCRH	UBIQUINOL-CYTOCHROME C REDUCTASE COMPLEX 11 KD PROTEIN PRECURSOR	R11698	Hs.73818
966	25499	UQCRH	UBIQUINOL-CYTOCHROME C REDUCTASE COMPLEX 11 KD PROTEIN PRECURSOR	R17676	Hs.73818
967	855843	UQCRB	UBIQUINOL-CYTOCHROME C REDUCTASE COMPLEX 14 KD PROTEIN	AA664284	Hs.131255
968	1160934	UQCRC2	UBIQUINOL-CYTOCHROME C REDUCTASE CORE PROTEIN 2 PRECURSOR	AA877491	
969	782800	UQCRFS1	UBIQUINOL-CYTOCHROME C REDUCTASE IRON-SULFUR SUBUNIT PRECURSOR	AA448184	Hs.3712
970	1955460	MAS2	UCR1_HUMAN P31930 UBIQUINOL- CYTOCHROME-C REDUCTASE COMPLEX CORE PROTEIN I	AI365986	
971	236034	UCP2	uncoupling protein 2 (mitochondrial, proton carrier)	H61243	Hs.80658
972	236034	UCP2	uncoupling protein 2 (mitochondrial, proton carrier)	H61242	Hs.80658
973	628529	EST	uncoupling protein 3 (mitochondrial, proton carrier)	AA192553	Hs.76640
974	628529	EST	uncoupling protein 3 (mitochondrial, proton carrier)	AA192136	Hs.76640
975	197932	UQCRB	UQCRB	R96352	Hs.77385
976	197932	UQCRB	UQCRB	R96395	Hs.77385

			1	1	1
0001	49464	UNG	uracil-DNA glycosylase	H15111	Hs.78853
977		UNG	uracil-DNA glycosylase	H15112	Hs.78853
978	49464	UROD	Uroporphyrinogen decarboxylase	AA424441	Hs.78601
979	760148	UROD	Uroporphyrinogen decarboxylase	AA424344	Hs.78601
980	760148	VEGF	vascular endothelial growth factor	R45059	
981	34778	VEGF	vascular endothelial growth factor	R19956	
982	34778		VDAC1	AI205945	
983	1762200	VDAC1	VDAC3	AI268057	
984	1911533	VDAC3	Voltage-dependent anion channel 1	AA044059	Hs.149155
985	486221	VDAC1	Voltage-dependent anion channel 1	AA044113	Hs.149155
986	486221	VDAC1	Voltage-dependent amon channel 2	AA857093	Hs.78902
987	1434908	VDAC2	Voltage-dependent amon charmel 2	AA460728	Hs.7381
988	796759	VDAC3	voltage-dependent anion channel 3	AA460900	Hs.7381
989	796759	VDAC3	voltage-dependent anion channel 3	AA400900	113.7501
	815683	WBSCR9	Williams-Beuren syndrome chromosome region 9	AA485132	Hs.194688
990	815683	WBSCR9	Williams-Beuren syndrome chromosome region 9	AA485131	Hs.194688
991	878676	XRCC5	X-ray repair complementing defective repair in Chinese hamster cells 5 (double- strand-break rejoining; Ku autoantigen, 80kD)	AA775355	Hs.84981
992		ZNF148	zinc finger protein 148 (pHZ-52)	N26148	
993	269381	ZNF9	zinc finger protein 9 (a cellular retroviral nucleic acid binding protein)	AA625995	Hs.2110
994	745503	ZNF9	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	<u> </u>	

Example 2 Mouse Mitochondrial Biology Array

A mouse mitochondrial biology array is made from clones representing expressed sequences. The clones placed on the array are shown in Table 4 which references sequence ID NOS:995-3040 provided herein setting forth the 5' and 3' sequences from these clones. See Tanaka, T.S. et al., (2000) "Genome-wide expression profiling of mid-gestation placenta and embryo using 15k mouse developmental cDNA microarray" *Proc. Natl. Acad. Sci. USA* 97:9127-9132. Equivalent clones useful as probes are listed in Table 5. The clones listed in Table 4 are preferable to the clones listed in Table 5. The clones identified in Table 4 are used to make a set of probes called Mouse Probe Set #2. The clones identified in Table 5 are used to make a set of probes called Mouse Probe Set #3. Control sequences are also placed this array. Controls include, but are not limited to blanks, DMSO, probes derived from plant sequences, sequence(s) not involved in mitochondrial biology, and poly adenine (40-60 nucleotides long). Sequences used to make probes for the mouse mitochondrial genes can also be found in GenBank Accession No. J01420, which provides the complete mouse mitochondrial genome. Preferably, the probes used for ATP8 and ATP6 do not cross-hybridize with each other.

Table

	abolism		n/Chromatin	n/Chromatin	n/Chromatin abolism abolism	n/Chromatin abolism abolism Stress	abolism abolism abolism	abolism abolism abolism abolism abolism
Mm.42805 Energy/Metabolism		47 Transcription/Chromatin		0135 Energy/Metabo	BG063066 Mm.100135 Energy/Metabolism BG076556 Mm.28696 Energy/Metabolism	Mm.128696 Energy/Metaboliss Mm.12970 Heat Shock/Stress	10135 Energy/Metabo 1696 Energy/Metabo 1970 Heat Shock/Str 1683 Energy/Metabo	Mm.12970 Heat Shock/Stress Mm.29683 Energy/Metabolism Mm.29683 Energy/Metabolism Mm.35389 Energy/Metabolism
Cluster ID Function	Mm.428		Mm.7447	Mm.744	Mm.744 Mm.100 Mm.286	Mm.1286 Mm.129	Mm.12970 Mm.29683 Mm.29683	Mm.744 Mm.286 Mm.286 Mm.296 Mm.353
Accession No.				BG063066	BG063066 BG076556	BG063066 BG076556 BG063078	BG076556 BG076556 BG063078	BG063066 BG076556 BG063078 BG063100
Clone description		Mus musculus major histocompatibility locus class III region:butyrophilin-like protein gene, partial cds; Notch4, PBX2, RAGE, lysophatidic acid acyl transferase-alpha, palmitoyl-protein	thioestera	thioestera Mus musculus heparan sulfate (glucosamine) 3-0-sulfotransferase 1 (Hs3st1), mRNA	thioestera Mus musculus heparan sulfate (glucosamine) 3-O-sulfotransferase 1 (Hs3st1), mRNA Rat farnesyltransferase beta subunit mRNA, complete cds	thioestera Mus musculus heparan sulfate (glucosamine) 3-0-sulfotransferase 1 (Hs3st1), mRNA Rat farnesyltransferase beta subunit mRNA, complete cds Mus musculus CPN10-like protein (Cpn10-rs1) gene, complete cds	thioestera Mus musculus heparan sulfate (glucosamine) 3-0-sulfotransferase 1 (Hs3st1), mRNA Rat farnesyltransferase beta subunit mRNA, complete cds Mus musculus CPN10-like protein (Cpn10-rs1) gene, complete cds Homo sapiens NADH dehydrogenase (ubiquinone) 1 beta subcomplex, 7 (18kD, B18) (NDUFB7), mRNA	thioestera Mus musculus heparan sulfate (glucosamine) 3-O-sulfotransferase 1 (Hs3st1), mRNA Rat farnesyltransferase beta subunit mRNA, complete cds Mus musculus CPN10-like protein (Cpn10-rs1) gene, complete cds Homo sapiens NADH dehydrogenase (ubiquinone) 1 beta subcomplex, 7 (18kD, B18) (NDUFB7), mRNA Mouse cytochrome c gene (MC1)
	805	us class III tein gene, RAGE, crase-alpha,		H3001C04 palmitoyl-protein thioestera this his paran sulfate (glucosamine) 3-O- (glucosamine) 3-O- (Hs. H3001F12 sulfotransferase 1 (Hs. 1), (Hs. 1)	e) 3-0-	QI OI		rotein thioestera fate (glucosamine) 3-0- rase 1 (Hs3st1), transferase beta subunit transferase beta subunit protein (Cpn10-rs1) gene ydrogenase (ubiquinone) 1 uplex, 7 (18kD, B18) c gene (MC1)
Clone No. Gene	H3001A07 Mm.42	reg Paa	H3001C04 pa	H3001C04 pal	H3001C04 pal he H3001F12 sul	H3001C04 pal H3001F12 sul H3001H04 Ra	H3001C04 pal H3001F12 sul H3001H04 Ra H3001H05 CF	H3001C04 pal hel H3001H04 Ra H3001H05 CF H3002A01 (N H3002B09 cy
SEQ ID NO:		966	1 L		0.7	0.1 0.5		

F0.	"Mus musculus RAB1, member RAS rran02E05 oncogene family (Rab1), mRNA"	Н	3G076435	Mm.14530	BG076435 Mm.14530 Signal Transduction
	oamide branched chain se E2 (Dbt),	Mus musculus dihydrolipoamide branched chain transacylase E2 (Dbt), mRNA	BG076436	Мт.3636	Energy/Metabolism
l H	"Mus musculus aurora-related kinase 2 Hannor 12 (ARK2) mRNA, complete cds"		BG076440	Mm.3488	Signal Transduction
	tic	Mus musculus glutamate-cysteine ligase catalytic subunit (GLCLC) mRNA, complete cds	BG076460 Mm.4368	Mm.4368	Energy/Metabolism
300	H3003D0SDHPS	Homo sapiens deoxyhypusine synthase (DHPS), mRNA	BG063127	Mm.28091	Mm.28091 Energy/Metabolism
30	H3003D08BAG2	Homo sapiens BCL2-associated athanogene 2 (BAG2), mRNA	BG076597	Mm.22449 Apoptosis	Apoptosis
03F0	H3003F02 ribonucleotide reductase	Mouse DNA for M2 subunit of mouse ribonucleotide reductase (EC 1.17.4.1)	BG076613 Mm.99	Мт.99	Energy/Metabolism
03F0	H3003F06 GluDH	Mus musculus glutamate dehydrogenase (Glud), mRNA	BG076616	Mm.10600	BG076616 Mm.10600 Energy/Metabolism
103F0	"Mus musculus calmodulin (Calm), H3003F07 mRNA"		BG076617	Mm.2648	Signal Transduction
03H("Mus musculus adenylate cyclase 7 H3003H01(Adcy7), mRNA"		BG076632	Mm.141400	BG076632 Mm.141400 Signal Transduction
 	H3004A02ISCU1 (ISCU), alternatively spliced	Homo sapiens ISCUI (ISCU) mRNA; complete cds, alternatively spliced	BG076641	Mm.29497	Mm.29497 Energy/Metabolism
1	mannosyl (alpha-1,3-)-glycoprotein beta-1,4-N-acetylglucosaminyltransferase,	Homo sapiens mannosyl (alpha-1,3-)-glycoprotein beta-1,4-N-acetylglucosaminyltransferase, isoenzyme B (MGAT4B), mRNA	BG076643	Mm.86759	BG076643 Mm.86759 Energy/Metabolism
3004A	H3004A04 isoenzyme B (MCA14B),	יייייין עביי אינטיוין ע טענענען טונען			

BG076647 Mm.100531 Energy/Metabolism	Signal Transduction		BG063219 Mm.153446 Signal Transduction	Signal Transduction	Mm.13145 Energy/Metabolism	Apoptosis	Energy/Metabolism	Protein Synthesis/Translational Control	Heat Shock/Stress
Mm.100531	Mm.1508	Mm.34776	Mm.153446	Mm.6355	Mm.13145	BG076705 Mm.21650 Apoptosis	Mm.27545		
BG076647	BG076653		BG063219	BG076689	BG076701	BG076705	BG076710	BG076718 Mm.12970	BG076720 Mm.2701
Mus musculus sarco(endo)plasmic reticulum calcium ATPase (SERCA2) gene, promoter region, exons 1-3, and partial cds					Mouse orotidine-5-monophosphate decarboxylase mRNA, 3 end	Mus musculus TF-1 apoptosis related protein 19 (Tfar19), mRNA	Mus musculus protein arginine N-methyltransferase 1 (Mrmt1) mRNA, complete cds, alternatively spliced	Mus musculus H19 and muscle-specific Nctol genes, complete sequence	
sarco(endo)plasmic reticulum calcium ATPase (SERCA2) gene, promoter region, exons 1-3, and partial cds	"Mouse mRNA for peripheral-type benzodiazepine receptor, complete cds"		"Mus musculus group V phospholipase H3004E03 A2 mRNA, complete cds"	"Mus musculus protein tyrosine H3004F07 phosphatase 4a2 (Pty4a2), mRNA"	orotidine-5-monophosphate H3004G09 decarboxylase, 3 end	TF-1 apoptosis related protein 19 (Tfar19),	protein arginine N-methyltransferase 1 (Mrnt1), alternatively spliced	muscle-specific Nctc1 genes,	"Mus musculus mammalian relative of H3005A06 DnaJ (Mrj-pending), mRNA"
sarco(e: ATPase H3004A09[region,	"Mouse H3004B06 benzodi	H3004C06	H3004E03	H3004F07	H3004G09	TF-1 ap H3004H01 (Tfar19)	protein H3004H06(Mmt1	H19 and 1	H3005A06
1034 1035	1036 1037	1038 1039	1040 1041	1042 1043	1044 1045	1046 1047	1048 1049	1050 1051	1052 1053

1054 1055	H3005B03	H3005B03 thioredoxin reductase 1 (Txnrd1),	Mus musculus thioredoxin reductase 1 (Txnrd1), mRNA	BG076728	Mm.44552	Mm.44552 Energy/Metabolism
1056 1057	H3005B06	H3005B06 amidophosphoribosyltransferase	Rattus norvegicus mRNA for amidophosphoribosyltransferase	BG076730	Mm.27743	Mm.27743 Energy/Metabolism
1058 1059	H3005D01 dipeptidyl	peptidase 4 (Dpp4),	Mus musculus dipeptidylpeptidase 4 (Dpp4), mRNA	BG076746	Mm.1151	Energy/Metabolism
1060	H3005D03	orotidine-5-monophosphate H3005D03 decarboxylase , 3 end	Mouse orotidine-5-monophosphate decarboxylase mRNA, 3 end	BG076748	Mm.802	Energy/Metabolism
1062 1063	"Homo s H3005E09 mRNA"	"Homo sapiens antiquitin 1 (ATQ1), mRNA"		BG076761	Mm.30250	Mm.30250 Signal Transduction
1064	H3005E10	ılus sodium/potassium ATPase unit	M.musculus mRNA for sodium/potassium. ATPase beta subunit BG076762.		Mm.4550	Energy/Metabolism
1066	"Mus mu H3005F06 kinase ml	"Mus musculus pelle-like protein kinase mRNA, complete cds"		BG076768	Mm.38241	Signal Transduction
1068	H3005G01	3-phosphate dehydrogenase	Mouse mRNA for glycerol-3-phosphate dehydrogenase, complete cds	BG076772	Mm.3711	Energy/Metabolism
1070	H3005G06		isomerase	BG076776 Mm.2412		Energy/Metabolism
1072 1073	H3005G12 methiony	1 tRNA synthetase	Homo sapiens mRNA for methionyl RNA synthetase, complete cds	BG063326	Mm.28173	Energy/Metabolism
1074 1075	H3006E03	H3006E03 cytochrome c-1 gene	mplete	BG076814	Mm.29196 Apoptosis	Apoptosis
1076 1077	H3006E12	UDP-glucose:glycoprotein H3006E12 glucosyltransferase 1 (HUGT1),	Homo sapiens UDP- glucose:glycoprotein glucosyltransferase 1 (HUGT1), mRNA BG076821	BG076821	Mm.45651	Mm.45651 Energy/Metabolism

olism	uction	uction	oolism	oolism	polism	bolism	luction	ural	bolism	bolism	bolism
BG076835 Mm.29824 Energy/Metabolism	Signal Transduction	Signal Transduction	Energy/Metabolism	Mm.29713 Energy/Metabolism	BG076882 Mm.10528 Energy/Metabolism	Energy/Metabolism	Signal Transduction	Matrix/Structural Proteins	Energy/Metabolism	Energy/Metabolism	BG063518 Mm.19185 Energy/Metabolism
Mm.29824	Mm.18041		Mm.426	Mm.29713	Mm.10528	Mm.28466	Mm.6698	Mm.7524	Mm.1776		Mm.19185
BG076835	BG076847	BG063409 Mm.3552	BG076872		BG076882	BG076887	BG076890	BG063514	BG063515		BG063518
Mus musculus puromycin-sensitive aminopeptidase (Psa), mRNA			Mus musculus glutathione S-transferase pi class (mGSTpiA) and (mGSTpiB) genes, complete cds		Mus musculus histidyl tRNA synthetase (Hars), mRNA	H.sapiens gene for 2-oxoglutarate carrier protein			Mus musculus ferritin heavy chain (Fth), mRNA		Mus musculus ATPase, H+ transporting, lysosomal I (Atp6i), mRNA
nycin-sensitive aminopeptidase	s musculus calmodulin synthesis	"Mus musculus circadian locomoter H3006H10 output cycles kaput (Clock), mRNA"	glutathione S-transferase pi class H3007E02 (mGSTpiA) and (mGSTpiB) genes	H3007G07\Mm.29713	histidyl fRNA synthetase (Hars),		Mus musculus GTP-binding protein (mSara) homologue mRNA, complete icds	Mus musculus dystroglycan 1 (DAG1)	H3008B06 ferritin heavy chain (Fth),	H3008B07 H3008B07	ATPase, H+ transporting, lysosomal I
puror H3006G03(Psa)	H3006H05	H3006H10	H3007E02	H3007G07	H3007G08histidy	H3008A02 protein	Mu (mS H3008A05cds	H3008B05	H3008B06	H3008B07	ATPase H3008B10(Amei)
1078	1080	1082	1084	1086	1088	1090	1092	1094	1096	1098	1100

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1102	"Mus mus protein PI	"Mus musculus PKCq-interacting protein PICOT (PICOT) mRNA,	pad	BG076907	Vm.29675	Mm.29675 Signal Transduction
	H3008C07	n-2,3-dioxygenase (TDO),	Mus musculus tryptophan-2,3- dioxygenase (TDO), mRNA	BG076911	Mm.21545	Mm.21545 Energy/Metabolism
	H3008C09	"HSP90=heat shock protein [mice, H3008C09]heart, mRNA Partial, 806 nt]"	7	AW538216 Mm.2180	1	Heat Shock/Stress
1108	H3008C12		Mus musculus adenylosuccinate lyase (Adsl), mRNA	BG076914	Мт.38151	Energy/Metabolism
1110	H3008D12H3008D1	H3008D12				Energy/Metabolism
			•			
1112	H3008E02	"Mus musculus caseinolytic protease, ATP-dependent, (E. coli) proteolytic H3008E02 subunit homolog (Cipp), mRNA"		BG076923	Mm.15243	Mm.15243 Signal Transduction
1114	H3008F09 Mm.3991	Mm.3991			Mm.3991	Energy/Metabolism
1116	,05800EH	s, alpha-D-galactosidase A ossomal protein (L44L), and vrosine kinase (Btk) genes	Mus musculus Btk locus, alpha-D-galactosidase A (Ags), ribosomal protein (L44L), and Brutons tyrosine kinase (Btk) genes, complete cds		Matrix/S Mm.30034 Proteins	Matrix/Structural Proteins
1118	M W W W W W W W W W W W W W W W W W W W	M.musculus gene cathepsin D, exons 6-	M.musculus gene cathepsin D, exons 6- M.musculus gene for cathepsin D, exons 6-9	AW538365		Apoptosis
1120	Н3008НО	arine palmitoyl transferase, subunit II ene; and unknown genes	Homo sapiens serine palmitoyl transferase, subunit II gene, complete cds, and unknown genes		Mm.29034	

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	BG063583 Mm.16757 Energy/Metabolism	Energy/Metabolism		Heat Shock/Stress	Mm.22514 Energy/Metabolism	Mm.14022 Energy/Metabolism	Signal Transduction	Mm.24293 Energy/Metabolism	Apoptosis	
	Mm.16757	Mm.2952	BG076966 Mm.41964	Mm.1843	Mm.22514	Mm.14022	Мт.8089	Mm.24293	Mm.105014	
	BG063583	BG063590	BG076966	BG063605 Mm.1843	BG076975	BG076988	BG063641		BG077079 Mm.105014Apoptosis	
	Mus musculus gibbon ape leukemia virus receptor (Slc20a1) gene, exon 10 and complete cds	Mus musculus flap structure specific endonuclease 1 (Fen1), mRNA	Mus musculus transglutaminase 1, K polypeptide (Tgn1), mRNA		Rattus norvegicus delta subunit of F1F0 ATPase gene, complete cds	Mus musculus cytochrome c oxidase, subunit VIIIa (Cox8a), mRNA		Homo sapiens mannosyl (alpha-1,6-)- glycoprotein beta-1,2-N- acetylglucosaminyltransferase (MGAT2), mRNA	Mus musculus tumor necrosis factor alpha converting enzyme (TACE) gene, exon 3, complete sequence	
	\Slc20a1	Fen1	Tgm1	"Mus musculus heat shock protein, 86 kDa 1 (Hsp86-1), mRNA"	H3009D02 ATPase delta	Cox8a	"Mus musculus lithium-sensitive myo- inositol monophosphatase A1 (IMPA1) mRNA, complete cds"	mannosyl (alpha-1,6-)-glycoprotein beta-1,2-N- acetylglucosaminyltransferase (MGAT2),	corosis factor alpha converting (TACE) gene, exon 3, sequence	Sociotocii
	H3009A09 SIc20a1	H3009B04 Fen1	H3009C03 Tgm1	"Mus r H3009C07kDa 1	H3009D02	H3009E08 Cox8a	"Mus m inositol H3009G03 mRNA,	mannosyl (beta-1,2-N acetylgluci H3010C12 (MGAT2),	tumor ne H3010D10 enzyme	904010ED 904010ED
	1122 1123	1124	1126 1127	1128 1129	1130 1131	1132 1133	1134 1135	1136 1137	1138 1139	1140

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	Apoptosis	Mm.25293 Energy/Metabolism	Signal Transduction	Energy/Metabolism			Signal Transduction		BG077186 Mm.34184 Heat Shock/Stress	Heat Shock/Stress	Energy/Metabolism	
	Mm.29816	Mm.25293	Mm.29842		Mm.2368	Mm.21743	Mm.57203		Mm.34184	Mm.4603	Mm.17362	
	BG077095 Mm.29816 Apoptosis	BG077121					BG077180		BG077186	BG077202	BG077218	
	Mus musculus programmed cell death 6 interacting protein (Pdcd6ip), mRNA	Mus musculus protein-L-isoaspartate (D-aspartate) O-methyltransferase 1 (Pcmt1), mRNA	H.sapiens mitogen inducible gene mig-2 2, complete CDS			Mus musculus Sid329 (Sid329), mRNA		Homo sapiens serine palmitoyl transferase, subunit II gene, complete ods: and unknown genes		4	Homo sapiens glyoxalase I (GLO1), mRNA	
	programmed cell death 6 interacting H3010F07 protein (Pdcd6ip),	rtate (D-aspartate) O- 1 (Pcmt1),	H3011B01 H.sapiens mitogen inducible gene mig-2	- H3011C08H3011C08	H3011E06 mTim17a	H3011E10 Sid329 (Sid329),	"House mouse mRNA for MAP kinase H3011H09kinase 3b, complete cds"	H3012 A 05 serine palmitoyl transferase	"Mus musculus heat shock 70 protein H3012A07 (Hsc70) gene, complete cds"	"Mus musculus scavenger receptor class B type I (mSR-BI) mRNA, complete scds"	GL01	Ill fire arriver or errecting the second
	H3010F07	H3011A03	H3011B01	- H3011C08	H3011E06	H3011E10	H3011H09	H3012A05	H3012A07	"Mu B ty H3012C06 cds"	H3012F03 GLO1	
,	1142 1143	1144	1146	1148	1150	1152	1154	1156	1158	1160	1162	1164

1166	H3012H0	"Mus musculus protein kinase Lkb1 H3012H04(Lkb1) and R29144/1 genes, partial cds"		BG063794	Mm.29947	BG063794 Mm.29947 Signal Transduction
1168 1169	H3013B02	M.musculus vacuolar adenosine H3013B02 triphosphatase subunit B	M.musculus mRNA for vacuolar adenosine triphosphatase subunit B	BG077252	Mm.10727	Mm.10727 Energy/Metabolism
1170	H3013B08	pyridoxal (pyridoxine, vitamin B6) H3013B08 kinase (PDXK),	Homo sapiens pyridoxal (pyridoxine, vitamin B6) kinase (PDXK), mRNA	BG063905	Mm.26103	Mm.26103 Energy/Metabolism
1171	H3013C0	H3013C04 "Mus musculus mRNA, complete cds"		AW539669 Mm.9846	Mm.9846	Heat Shock/Stress
1173 1174	H3013D03	"Mus musculus RAB10, member RAS H3013D03oncogene family (Rab10), mRNA"		BG077272	Мт.21985	Signal Transduction
1175 1176	H3013D11	H3013D11 Mouse metallothionein II (MT-II) gene		BG063925	Mm.89170	Mm.89170 Heat Shock/Stress
1177	adenyl; H3013E04 (CAP),	yl cyclase-associated protein	Homo sapiens adenylyl cyclase- associated protein (CAP), mRNA		Mm.43660	
1179	dolichyl-; acetylglu H3013E07 (Dpagt2),	phosphate alpha-N- cosaminephosphotransferase 2	Mus musculus dolichyl-phosphate alpha-N- acetylglucosaminephosphotransferase 2 (Dpagt2), mRNA	BG063933	Mm.18353	Energy/Metabolism
1181 1182	H3013F01	nolevulinate dehydratase (Lv),	olevulinate	BG063937		Energy/Metabolism
1183 1184	H3013F02	"Mus musculus aurora-related kinase 1 H3013F02 (ARK1) mRNA, complete cds"		BG077290	Mm.11738	BG077290 Mm.11738 Signal Transduction
1185 1186	survivin4 survivin1 H3013G10products	10, survivin121, and 140 genes, alternative splice	Mus musculus survivin40, survivin121, and survivin140 genes, alternative splice products, complete cds	BG077309	Mm.8552	Apoptosis

1187	H3014A05	H3014A05 partial aminopeptidase B (APB gene)	Homo sapiens partial mRNA for aminopeptidase B (APB gene)	BG077324 P	Mm.29706	Mm.29706 Energy/Metabolism
	"Rattus 1 with sme	"Rattus norvegicus protein associating with small stress protein PASSI (Pass1)			Mm.45272	Mm.45272 Heat Shock/Stress
1189	H3014A00	tRNA-ribosyltransferase 507).	Mus musculus queuine tRNA- ribosyltransferase (LOC60507), mRNA BG077329		Mm.24178	Energy/Metabolism
1192	H3014B06				Mm.29849	Energy/Metabolism
1194	Mus mi	Mus musculus surfeit gene 4 (Surf4), 8 mRNA		BG063996	Mm.2795	
1196	H3014C10	0				Heat Shock/Stress
1198	H3014C12	solute carrier family 28 (sodium- coupled nucleoside transporter), member 2 (Slo28a2),	Mus musculus solute carrier family 28 (sodium-coupled nucleoside transporter), member 2 (Slc28a2), mRNA	BG077361	Mm,29510	Mm.29510 Energy/Metabolism
1200	H3014E00	UDP-Gal:betaGlcNAc beta 1,3- galactosyltransferase, polypeptide 4 H3014E06 (B3galt4),	Mus musculus UDP-Gal:betaGlcNAc beta 1,3-galactosyltransferase, polypeptide 4 (B3galt4), mRNA	BG077355	Mm.11132	BG077355 Mm.11132 Energy/Metabolism
1202	H3014E0	H3014E07 signal peptidase 21kDa subunit	Rattus norvegicus mRNA for signal peptidase 21kDa subunit, complete cds		Mm.27800	BG077356 Mm.27800 Energy/Metabolism
1204	H3014F0:	"Mus musculus S100 calcium-binding H3014F05 protein A13 (S100a13), mRNA"			Mm.6523	Signal Transduction
1205 1206	H3014F0	UDP-Gal:betaGlcNAc beta 1,4- galactosyltransferase, polypeptide 1 H3014F06 (B4galt1),	Mus musculus UDP-Gal:betaGlcNAc beta 1,4- galactosyltransferase, polypeptide 1 (B4galt1), mRNA	BG077374	Mm.15622	BG077374 Mm.15622 Energy/Metabolism

1207 1208	phosphc H3014G07115 (Pea	phosphoprotein enriched in astrocytes 15 (Peal 5),	Mus musculus phosphoprotein enriched in astrocytes 15 (Pea15), mRNA	BG064035	Mm.544	Apoptosis
1209 1210	protease H3014G08 subunit,	protease (prosome, macropain) 26S subunit, ATPase 5 (Psmc5),	Mus musculus protease (prosome, macropain) 26S subunit, ATPase 5 (Psmc5), mRNA	BG077385	Mm.665	Energy/Metabolism
1211 1212	protease H3014G09 subunit,	protease (prosome, macropain) 26S subunit, ATPase 5 (Psmc5),	Mus musculus protease (prosome, macropain) 26S subunit, ATPase 5 (Psmc5), mRNA		Mm.665	Energy/Metabolism
1213 1214	H3014H07	H3014H07 tyrosyl-tRNA synthetase (YARS),	Homo sapiens tyrosyl-tRNA synthetase (YARS), mRNA	BG077395	Mm.20353	Mm.20353 Energy/Metabolism
1215 1216	H3015A01	major histocompatibility locus class III region:butyrophilin-like protein gene, partial cds; Notch4, PBX2, RAGE, lysophatidic acid acyl transferase-alpha, H3015A01 palmitoyl-protein thioestera	Mus musculus major histocompatibility locus class III region:butyrophilin-like protein gene, partial cds; Notch4, PBX2, RAGE, lysophatidic acid acyl transferase-alpha, palmitoyl-protein thioestera		Mm.27416	
121 <i>7</i> 1218	H3015B03	hypothetical protein FLJ10535 H3015B03 (FLJ10535),	Homo sapiens hypothetical protein FLJ10535 (FLJ10535), mRNA	BG077413	Mm.24187	Mm.24187 Energy/Metabolism
1219 1220	methion H3015C02 subunit	ine adenosyltransferase alpha gene fragment	Homo sapiens methionine adenosyltransferase alpha subunit gene fragment	BG064075	Mm.29815	Mm.29815 Energy/Metabolism
1221 1222	H3015D05 MSTP029		Homo sapiens MSTP029 mRNA, complete cds	BG077438	Mm.30147	BG077438 Mm.30147 Energy/Metabolism
1223 1224	H3015E06	Mus musculus Tuf receptor-associated factor 1 (Trafl), factor 1 (Trafl), mRNA	otor-associated	BG064103	Mm.12898 Apoptosis	Apoptosis
1225 1226	H3016C10					Heat Shock/Stress

				-		
1227 1228	H3016D01 Mm.2020	Mm.20201			Mm.20201	Energy/Metabolism
1229 1230	H3016D08	novirus E1B 19 kDa- g protein 3-like (Bnip31),	Mus musculus BCL2/adenovirus E1B 19 kDa-interacting protein 3-like (Bnip31), mRNA	BG077518	Mm.29820 Apoptosis	Apoptosis
1231	Mus GTJ H3016E10 cds	Mus musculus mG28K mRNA for GTP-binding protein like 1, complete cds		BG077528	Mm.28954	
1233	H3016F03 Tim44	44	Rattus norvegicus mRNA for Tim44, complete cds		Mm.34791	Matrix/Structural Proteins
1235 1236	H3016F07 M.muscu	lus glutamyl-tRNA synthetase	is mRNA for glutamyl-tRNA	BG064194	Mm.27190	Mm.27190 Energy/Metabolism
1237	H3016F11					Heat Shock/Stress
1238 1239	ornithir H3017A02(Oaz2),	ne decarboxylase antizyme 2	Mus musculus ornithine decarboxylase antizyme 2 (Oaz2), mRNA	BG077570 Mm.675	Mm.675	Energy/Metabolism
1240 1241	"Mus mu nucleotic H3017C11 pending)	"Mus musculus RAN guanine nucleotide release factor (Rangnrf- pending), mRNA"		BG077598	Mm.143774	Mm.143774 Signal Transduction
1242 1243	"Homo s E6-AP (domain s H3017D04(RLD) 1	"Homo sapiens hect (homologous to the E6-AP (UBE3A) carboxyl terminus) domain and RCCI (CHCI)-like domain (RLD) I (HERCI), mRNA"		BG064250	Mm.102717	BG064250 Mm.102717 Heat Shock/Stress
1244 1245	"Mus mu H3017E04 (Ctsz) m	"Mus musculus cathepsin Z precursor (Ctsz) mRNA, complete cds"		BG077611	Mm.115	Signal Transduction
1246 1247	"Mus mu H3017E11 protein 7	"Mus musculus retinoblastoma binding protein 7 (Rbbp7), mRNA"	1	AW544081 Mm.1603		Heat Shock/Stress

1248 1249	phenyla H3017G06 subunit	phenylalanyl tRNA synthetase beta subunit (Frsb)	Mus musculus phenylalanyl tRNA synthetase beta subunit (Frsb) mRNA, complete cds	BG077633	Mm.28922	BG077633 Mm.28922 Energy/Metabolism
1250 1251	H3018A02B4galt3	B4galt3	Mus musculus beta-1,4- galactosyltransferase III (B4galt3), mRNA	BG077551	Mm.150720	Mm.150720 Energy/Metabolism
1252 1253	H3018A04 APB	APB	Homo sapiens partial mRNA for aminopeptidase B (APB gene)	BG077553		Energy/Metabolism
1254 1255	"Mus m chapero H3018A08 subunit,	"Mus musculus Cctq gene for chaperonin containing TCP-1 theta subunit, complete cds"		BG064304	Мт.17989	Mm.17989 Heat Shock/Stress
1256 1257	H3018A11 Cyp17	Cyp17	Mus musculus cytochrome P450, 17 (Cyp17), mRNA	BG064307	Mm.1262	Energy/Metabolism
1258 1259	H3018B08	H3018B08 thimet oligopeptidase	Mus musculus thimet oligopeptidase mRNA, complete cds	BG077656	Mm.26995	
1260 1261	H3018F06			·		Heat Shock/Stress
1262 1263	H3018G08 UDP-gal	actose 4 epimerase (GALE)	Homo sapiens UDP-galactose 4 epimerase (GALE) gene, complete cds	BG077710	Мт.20363	BG077710 Mm.20363 Energy/Metabolism
1264	H3019A07	ATP binding protein associated with H3019A07 (cell differentiation (APACD),	Homo sapiens ATP binding protein associated with cell differentiation (APACD), mRNA		Mm.28438	Mm.28438 Energy/Metabolism
1265 1266	ATPase, H3019B01 polypept	Na+/K+ transporting, beta 1 ide (Atp1b1),	Mus musculus ATPase, Na+/K+ transporting, beta 1 polypeptide (Atp1b1), mRNA	BG077733 Mm.4550	Mm.4550	Energy/Metabolism
1267 1268	H3019B03	acyl-coenzyme A:cholesterol acyltransferase [mice, peritoneal H3019B03 macrophages, , 3041 nt]	acyl-coenzyme A:cholesterol acyltransferase [mice, peritoneal macrophages, mRNA, 3041 nt]	BG077735	Matrix/S Mm.28099 Proteins	Matrix/Structural Proteins

1269	Mus musor	Mus musculus microtubule-associated protein 4 (Mtap4), mRNA		Matrix/S BG077742 Mm.12625 Proteins	Mm.12625	Matrix/Structural Proteins
1271	H3019C06	rotein (LOC51012),	Homo sapiens CGI-107 protein (LOC51012), mRNA		Mm.30065	
			Mus musculus core1 UDP-galactose:N-			
1273		eta 1,3-		CAPPROO	1007E0	
1274	H3019C11	galactosyltransferase (Clgalt1)	complete cds	•	7017017111111	(G)
1275	"Mu" (mSi	"Mus musculus GTP-binding protein (mSara) homologue mRNA, complete		BG077753	Mm.6698	Signal Transduction
1277		"Mus musculus protein kinase C inhibitor (mPKCI) mRNA, complete		035550000	10 435	Simel Transduction
1278	H3019D09 cds"	cds"		867//29	C74-TIIIVI	Olgial Hansdavion
1279	H3019E05	H3019E05 Mm.38994			Mm.38994	Energy/Metabolism
1281			Homo sapiens F1F0-type ATP synthase	BG077769	Mm 102755	Mm 102755 Energy/Metabolism
1282	H3019E12	H3019E12 F1F0-type ATP synthase subunit d	subunit a meny, compiete cas		201-01-01-01-01-01-01-01-01-01-01-01-01-0	(a)
1283 1284	H3019F05	H3019F05 Mouse tyrosine kinase (c-abl) mRNA		BG077774 Mm.1318	Mm.1318	Signal Transduction
1285		domesticus tumor necrosis factor	Mus musculus domesticus mRNA for tumor necrosis factor receptor p60			
1286	H3019F06 gene)	gene)	homologue 1 (Tnfrh1 gene)	BG077775	Mm.103353 Apoptosis	Apoptosis
1287 1288	H3019F07	"Mus musculus serine/threonine kinase H3019F07 10 (Stk10), mRNA"		BG077776 Mm.8235	Mm.8235	Signal Transduction
1289 1290	H3019F10	H3019F10 phosphomannomutase 2 (Pmm2),				Energy/Metabolism

1291 1292	H3019G05	H3019G05 alkaline phosphatase 2, liver (Akp2),	Mus musculus phosphomannomutase 2 (Pmm2), mRNA	BG077784 Mm.9699	Mm.9699	Energy/Metabolism
1293 1294	ATPas H3019H07(Atpl),	ATPase-like vacuolar proton channel (Atpl),	Mus musculus alkaline phosphatase 2, iver (Akp2), mRNA	BG077796	Mm.1265	Energy/Metabolism
1295 1296	H3019H11	H3019H11 H3019F10	Mus musculus ATPase-like vacuolar proton channel (Atpl), mRNA	BG077799	Mm.30155	Mm.30155 Energy/Metabolism
1297 1298	nucle H3020A02[gene)	oside diphosphatase (ER-UDPase	Mus musculus mRNA for nucleoside diphosphatase (ER-UDPase gene)	BG077800	Mm.10211	Mm.10211 Energy/Metabolism
1299 1300	serine pa H3020A07/gene; an	ılmitoyl transferase, subunit II d unknown genes	Homo sapiens serine palmitoyl transferase, subunit II gene, complete cds; and unknown genes		Mm.22626	
1301 1302	H3020B08 ferritin	ferritin light chain 1 (Ft11),	Mus musculus ferritin light chain 1 (Ft11), mRNA	BG077812	Mm.7500	Energy/Metabolism
1303 1304	"Mouse H3020C02 5' end"	"Mouse metallothionein-I (MT-I) gene, 5' end"		BG077818	Mm.2041	Heat Shock/Stress
130 5 1306	isocitrat H3020D10 alpha (II	e dehydrogenase 3 (NAD+) NH3A),	Homo sapiens isocitrate dehydrogenase 3 (NAD+) alpha (IDH3A), mRNA	BG077913	Mm.29051	Mm.29051 Energy/Metabolism
1307 1308	"Mus m kDa heat H3020E01 complete	"Mus musculus wagneri gene for 105- kDa heat shock protein, exon 18 and complete cds"		BG064500	Mm.34828	Mm.34828 Heat Shock/Stress
1309 1310	H3020H04	H3020H04 squalene epoxidase (Sqle),	Mus musculus squalene epoxidase (Sqle), mRNA	BG077950	Мт.22663	BG077950 Mm.22663 Energy/Metabolism
1311 1312	BALB/c H3020H07 (GDI-1)	: GDP-dissociation inhibitor , partial cds	Mus musculus BALB/c GDP- dissociation inhibitor (GDI-1) mRNA, partial cds	BG077953	Mm.28084	BG077953 Mm.28084 Energy/Metabolism

1313			Mouse man 6-P receptor (46MPR) mRNA. complete cds	BG064540	Mm.1358	Energy/Metabolism
1314	H3020H10 man 0-F 15	ed cell death 6 (Pdcd6),	omed cell death 6	BG077957	Mm.24254 Apoptosis	Apoptosis
1317	H3021A11	finger-like protein (Tim13)	mall zinc finger-like mRNA, complete cds	BG077964	Mm.22201	Mm.22201 Energy/Metabolism
1319	H3021A12	us (clone S5) WRS tRNA ligase	M.musculus (clone S5) WRS mRNA for hyptophantRNA ligase	BG077965	Mm.38433	Energy/Metabolism
1321	H3021B05	ne	M.musculus CI-23 kD gene	BG077969	Mm.44227	Energy/Metabolism
1323	H3021B08	mitochondrial carrier peroxisomal membrane protein, 34 kDa member H3021B0817 PMP34			Mm.306	
1325	H3021C09	н3021С09 Н3021С09				Apoptosis
1327	H3021D07	"Mus musculus proliferin related protein		BG077825	Mm.3258	Signal Transduction
1329	H3021E11	sine triphosphatase	Mus musculus vacuolar adenosine triphosphatase subunit A gene, complete cds	BG064589	Mm.29771	Mm.29771 Energy/Metabolism
1331	H3021F01	aspartyl a	Mus musculus aspartyl aminopeptidase (Dnpep), mRNA	BG077839	Mm.24680	Mm.24680 Energy/Metabolism
1333	H3021F08	"Mus musculus chaperonin subunit 7 H3021F08 (eta) (Cct7), mRNA."		BG077843	Mm.914	Heat Shock/Stress
1335	H3021G03	"Mus musculus serine/threonine protein phosphatase type 1 alpha mRNA, H3021G03 complete cds"		BG077848	BG077848 Mm.1970	Signal Transduction

Signal Transduction	Energy/Metabolism	Signal Transduction	Energy/Metabolism	Mm.27890 Energy/Metabolism	Energy/Metabolism	Energy/Metabolism	Mm.29010 Energy/Metabolism	Mm.102791Energy/Metabolism	Mm.29998 Energy/Metabolism	Energy/Metabolism
	Mm.28864	Mm.40644	Mm.41801	Mm.27890	Мт.2820	Mm.3317	Mm.29010	Mm.102791	Mm.29998	/
BG077860 Mm.905	BG064623	BG077872	BG064641	BG064654	BG064655	BG077995	BG064664	BG078006	BG064680	BG078023 Mm.2462
	Cricetulus griseus mRNA for Phosphatidylglycerophosphate synthase, complete cds		Homo sapiens ribonuclease P (14kD) (RPP14), mRNA	Homo sapiens dolichyl-phosphate beta- glucosyltransferase mRNA, complete cds	Mus musculus galactokinase gene, complete cds	Mouse beta-glucuronidase gene, complete cds	Mus musculus GLUT4 vesicle protein mRNA, partial cds	Homo sapiens a disintegrin-like and metalloprotease (reprolysin type) with thrombospondin type 1 motif, 4 (ADAMTS4), mRNA	Mus musculus malic enzyme, supernatant (Mod1), mRNA	Mus musculus 26S protease ATPase (mss1) mRNA, partial cds
"Mus musculus Ras suppressor protein 1 (Rsu1), mRNA"	Cricetulus griseus Cricetulus griseus H3022A04Phosphatidylglycerophosphate synthase complete eds	"Mus musculus RAB23, member RAS H3022A05 oncogene family (Rab23), mRNA"			H3022D03 galactokinase gene	gene	GLUT4 vesicle protein , partial cds	a disintegrin-like and metalloprotease (reprolysin type) with thrombospondin H3022E09 type 1 motif, 4 (ADAMTS4),		al
"Mus m H3021H03 (Rsu1),	H3022A04	H3022A05	H3022B12	dolichyl H3022D02 glucosy	H3022D03	H3022D09	H3022E01 GLUT4	H3022E09	H3022F06	26S H3022G02 cds
1337 1338	1339 1340	1341	1343 1344	1345 1346	1347 1348	1349 1350	1351 1352	1353 1354	1355 1356	13 <i>57</i> 1358

1359	26S	protease ATPase (mss1) , partial	Mus musculus 26S protease ATPase (mss1) mRNA, partial cds	BG078024 Mm.2462		Energy/Metabolism
	serine/thr	ne/threonine kinase 17b (apoptosis- ucine) (STK17B),	Homo sapiens serine/threonine kinase 17b (apoptosis-inducing) (STK17B), mRNA	BG078034 N	Mm.25559 Apoptosis	Apoptosis
			д	BG064691	Mm.849	Signal Transduction
1364	H3022H00	ursor		RG064717	Мт.3830	Energy/Metabolism
1366	H3023A10 (Fpgs)	O (FPES)			1 1	Heat Shock/Stress
1368	H3023B01					
1369	H3023B1	"Mouse mRNA for HSP60 protein H3023B11(clones 3T3-7,-9, and -M1)"		BG064728 Mm.1777	·	Heat Shock/Stress
1371	T200201	Transable Manies child for heat shock protein 65		BG064729	Mm.1777	Heat Shock/Stress
13/2	19270CH	A INTOLOGO COLOR SERVICIO COLOR SERV				
1373	H3023C0	Homo sapiens nuclear receptor outung H3023C05 protein (NRBP), mRNA		BG064733	Mm.22029	
1375	H3023C0	H3023C07 peroxiredoxin 5 (Prdx5),	Mus musculus peroxiredoxin 5 (Prdx5), mRNA	BG064735 Mm.6587	Mm.6587	Energy/Metabolism
1377	H3023C0	H3023C08 ribonucleotide reductase M1 (Rrm1),	Mus musculus ribonucleotide reductase M1 (Rrm1), mRNA	BG064736	Mm.656	Energy/Metabolism
1379	H3023C0	H3023C09\Mm.29735			Mm.29735	Mm.29735 Energy/Metabolism
2001				,		
1381 1382	H3023D(Mus musculus phosphoglycerate kinase H3023D06(Pgk1-ps1) processed pseudogene		BG064745	Mm.188	Signal Transduction

Energy/Metabolism	BG078081 Mm.100638 Energy/Metabolism	Energy/Metabolism	Signal Transduction	Heat Shock/Stress	Heat Shock/Stress	Mm.22374 Energy/Metabolism	Heat Shock/Stress	Heat Shock/Stress	Heat Shock/Stress
Mm.30130	Mm.100638	Mm.10406	Mm.3191	Mm.1813	Mm.1813	Mm.22374	Mm.1843	Mm.1843	Mm.1843
BG064747	BG078081	BG078095	BG064765	BG064769	BG064770	BG064771	BG064772	BG064774	BG064775
Human glutamate pyruvate transaminase (GPT) gene, complete cds BG064747	Mus musculus partial mRNA for xylosyltransferase II (XT-II gene)	Homo sapiens SDHD gene for small subunit of cytochrome b of succinate dehydrogenase, complete cds				Mus musculus glucosamine-6- phosphate deaminase (Gnpda) mRNA, complete cds			
mate pyruvate transaminase (GPT)	l xylosyltransferase II (XT-II	SDHD gene small subunit of cytochrome b of succinate H3023E11 dehydrogenase	"Mus musculus phosphatidylinositol-4- phosphate 5-kinase, type 1 beta H3023F02 (Pip5k1b), mRNA"	"Mus musculus Ccte gene for chaperonin containing TCP-1 epsilon subunit, complete cds"	"Mus musculus Ccte gene for chaperonin containing TCP-1 epsilon subunit, complete cds"	nine-6-phosphate deaminase	"Mus musculus heat shock protein, 86 kDa 1 (Hsp86-1), mRNA"	"Mus musculus heat shock protein, 86 kDa 1 (Hsp86-1), mRNA"	"Mus musculus heat shock protein, 86 H3023G01]kDa 1 (Hsp86-1), mRNA"
gluta H3023D08 gene	partia H3023D09 gene)	H3023E11	H3023F02	"Mus m chapero H3023F07 subunit,	"Mus m chapero H3023F08 subunit,	glucosar H3023F09 (Gnpda)	"Mus H3023F10 kDa 1	"Mus 1 H3023F12 kDa 1	H3023G01
1383 1384	1385 1386	1387 1388	1389 1390	1391 1392	1393 1394	1395 1396	1397 1398	1399 1400	1401 1402

		phosphate	Mus musculus phosphoribosyl pyrophosphate synthetase 1 (Prps1), mRNA	BG064779	Mm.25198	Energy/Metabolism
	H30Z3G05	1 (ribsi),	sculus ferritin L-subunit gene 4, complete cds	BG064794	Mm.7500	Energy/Metabolism
	H3023H09 territio L.	subtuil gene exons 1-4	subunit gene	BG064795	Mm.7500	Energy/Metabolism
1407 1408 1409	H3023H10 ferritin L	subunit gene exons 1-4	subunit gene	BG064796	Mm.7500	Energy/Metabolism
1410	lactate (H3023H12(Tdh1).	lactate dehydrogenase 1, A chain (f.dh.1).	Mus musculus lactate dehydrogenase 1, A chain (Ldh1), mRNA	BG064797	Mm.26504	Energy/Metabolism
1412 1413	H3024A02	H3024A02 polymerase, gamma (Polg),	Mus musculus polymerase, gamma (Polg), mRNA	BG064799	Mm.3616	DNA Replication
1414 1415	H3024A06	H3024A06 alcohol dehydrogenase 5 (Adh5),	Mus musculus alcohol dehydrogenase 5 (Adh5), mRNA	BG064803	Mm.3874	Energy/Metabolism
1416	H3024A1	DNA M2 subunit of ribonucleotide H3024A11 reductase (EC 1.17.4.1)	Mouse DNA for M2 subunit of mouse ribonucleotide reductase (EC 1.17.4.1)	BG078138	Мт.99	Energy/Metabolism
1418	H3024B0	"Mus musculus chaperonin subunit 3 H3024B03(gamma) (Cct3), mRNA"		BG064811	Mm.3576	Heat Shock/Stress
1420	H3024B0			BG064812	Mm.3576	Heat Shock/Stress
1422 1423	H3024B0	"Mus musculus chaperonin subunit 3 H3024B05 (gamma) (Cct3), mRNA"		BG064813	Mm.3576	Heat Shock/Stress
1424 1425	H3024B0	"Mus musculus chaperonin subunit 3 H3024B06 (gamma) (Cct3), mRNA"		BG064814	Mm.3576	Heat Shock/Stress
1426 1427	H3024B12	2				Heat Shock/Stress

1428 1429	H3024C02	"Homo sapiens oxidative-stress H3024C02 responsive 1 (OSR1), mRNA"		BG078151	Mm.52786	Mm.52786 Heat Shock/Stress
1430 1431	H3024C04	H3024C04 phosphoglycerate mutase 1 (Pgam1)	Mus musculus phosphoglycerate mutase 1 (Pgam1) mRNA, complete cds BG064823	i	Mm.16783	Energy/Metabolism
1432 1433	H3024C07 HEXA	HEXA gene, exons 2 - 14	M.musculus HEXA gene, exons 2 - 14	BG064825	Mm.2284	Energy/Metabolism
1434 1435	"Mus m H3024C11 kDa 1 ("Mus musculus heat shock protein, 86 kDa 1 (Hsp86-1), mRNA"		BG064829	Mm.1843	Heat Shock/Stress
1436 1437	H3024E08	H3024E08 direct IAP binding protein with low PI	Mus musculus direct IAP binding protein with low PI mRNA, complete cds	BG064850	Mm.46716 Apoptosis	Apoptosis
1438 1439	H3024F05	Mus musculus putative CCAAT binding factor 1 (mCBF) mRNA, alternatively H3024F05 spliced transcript mCBF1, complete cds		BG064857	Mm.24169	Transcription/Chromatin
1440 1441	serine hy H3024F06 (soluble)	serine hydroxymethyl transferase 1 (soluble) (Shmt1),	Mus musculus serine hydroxymethyl transferase 1 (soluble) (Shmt1), mRNA	BG078187	Mm.45993	Energy/Metabolism
1442 1443	H3024F07	H3024F07 fumarate hydratase (FH),	Homo sapiens fumarate hydratase (FH), mRNA	BG064859	Mm.41502	Energy/Metabolism
1444 1445	phosphol H3024G02 synthetas	ribosyl pyrophosphate se I (Prps1),	Mus musculus phosphoribosyi pyrophosphate synthetase 1 (Prps1), mRNA	BG064866	Mm.25198	Mm.25198 Energy/Metabolism
1446 1447	H3024G03	H3024G03 K+ channel, sequence	Rattus norvegicus K+ channel mRNA, . sequence	BG078196	Mm.40482	Mm.40482 Energy/Metabolism
1448 1449	H3024G07 Murine I	Murine L-myc gene		BG064871	Mm.1055	Signal Transduction
1450 1451	"Mus mu H3024H02 mRNA, 1	"Mus musculus p53 binding protein 1 mRNA, partial cds"		BG078205	Mm.25231	Mm.25231 Signal Transduction

1452	H3024H12	"Mus musculus breast heat shock 73 H3024H12brotein (hsc73) mRNA, complete cds"	·	BG064886 Mm.2944	1	Heat Shock/Stress
	"Mus mr responsi H3025A01 mRNA"	"Mus musculus interferon alpha responsive protein (15 kDa) (Ifig15), mRNA"		BG078215	Мт.21761	Heat Shock/Stress
	H3025C05	ehydrogenase, soluble (Mor2),	Mus musculus malate dehydrogenase, soluble (Mor2), mRNA	BG064914	Mm.3156	Energy/Metabolism
1458 1459	H3025D10	Aga=aspartylglucosaminidase [mice, H3025D10 liver, brain, Partial, 1191 nt]	Aga=aspartylglucosaminidase [mice, liver, brain, mRNA Partial, 1191 nt]	BG064929	Mm.41591	Energy/Metabolism
1460 1461	H3025E07	1Se	Mus musculus ADP-ribosylarginine hydrolase mRNA, complete cds	BG064935	Mm.20047	Mm.20047 Energy/Metabolism
1462 1463	H3025F10	succinate dehydrogenase Ip subunit,	Mus musculus succinate dehydrogenase Ip subunit mRNA, partial cds	BG064949	Mm.29141	Mm.29141 Energy/Metabolism
14641465	14641465 H3026B03 HSPC14;	5 protein (HSPC145),	Homo sapiens HSPC145 protein (HSPC145), mRNA		Mm.29904	
1466	H3026B04	ibosyl pyrophosphate nsferase (PPAT),	Homo sapiens phosphoribosyl pyrophosphate amidotransferase (PPAT), mRNA	BG064988	Mm.27743	Mm.27743 Energy/Metabolism
1468	H3026B06				Mm.100588	Mm.100588 Energy/Metabolism
1470 1471	H3026B07	H3026B07 ADP-ribosylation factor 1 (Arf1),	Mus musculus ADP-ribosylation factor 1 (Arf1), mRNA	BG078294	Mm.6836	Energy/Metabolism
1472 1473	H3026B11		Mus musculus ATPase, Cu++ transporting, beta polypeptide (Atp7b), mRNA	BG078297	Mm.102506	BG078297 Mm.102506 Energy/Metabolism

Energy/Metabolism	Heat Shock/Stress	Energy/Metabolism	Energy/Metabolism	BG065033 Mm.18759 Energy/Metabolism	Energy/Metabolism	Heat Shock/Stress	Heat Shock/Stress	Mm.34830 Energy/Metabolism	Wm.29815 Energy/Metabolism
Mm.3308	Mm.27897	BG078316 Mm.12983	Mm.1090	Mm.18759	Mm.29856	Мт.914	Mm.914	Мт.34830	Mm.29815
BG065012 Mm.3308	BG065015	BG078316	BG065030	BG065033	BG078326	BG065035	BG065036	BG078342	BG065061
Mus musculus tyrosine 3- monooxygenasc/tryptophan 5- monooxygenase activation protein, eta polypeptide (Ywhah), mRNA		Homo sapiens cDNA FLJ12225 fis, clone MAMMA1001139, weakly similar to SRE-2 PROTEIN	M.musculus GSHPx gene	Mus musculus acetyl-Coenzyme A dehydrogenase, short chain (Acads), mRNA	H.sapiens mRNA for phosphoenolpyruvate carboxykinase			Homo sapiens mannose phosphate isomerase (MPI), mRNA	Homo sapiens methionine adenosyltransferase alpha subunit gene fragment
Mus musculus tyrosine 3- tyrosine 3-monooxygenase/tryptophan monooxygenase/tryptophan 5- 5-monooxygenase activation protein, eta monooxygenase activation protein, eta polypeptide (Ywhah), mRNA	"Mus musculus heat shock protein, H3026D10DNAJ-like 2 (Hsj2), mRNA"	cDNA FLJ12225 fis, clone MAMMA1001139, weakly similar to H3026E07 SRE-2 PROTEIN	H3026F02 M.musculus GSHPx gene	acetyl-Coenzyme A dehydrogenase, H3026F05 short chain (Acads),	H.sapiens phosphoenolpyruvate	"Mus musculus Ccth gene for chaperonin containing TCP-1 eta H3026F07 subunit, complete eds"	"Mus musculus Ccth gene for chaperonin containing TCP-1 eta subunit, complete cds"	H3026G10 mannose phosphate isomerase (MPI),	methionine adenosyltransferase alpha subunit gene fragment
H3026D06	H3026D10	H3026E07	H3026F02	H3026F05	H3026F06	H3026F07	H3026F08	H3026G10	methior H3026H09 subunit
1474 1475	1476	1478	1480 1481	1482 1483	1484 1485	1486 1487	1488 1489	1490 1491	1492 1493

1494 1495	puron H3026H11(Psa),	nycin-sensitive aminopeptidase	Mus musculus puromycin-sensitive aminopeptidase (Psa), mRNA	BG065063	Mm.29824	BG065063 Mm.29824 Energy/Metabolism
1496	ATPa H3027A10(Atpl)	se-like vacuolar proton channel	Mus musculus ATPase-like vacuolar proton channel (Atpl), mRNA	BG065073	Mm.30155	Mm.30155 Energy/Metabolism
1498 1499	"Homo sa H3027B02 protein 4	"Homo sapiens SH3-domain binding protein 4 (SH3BP4), mRNA"		BG078369	Mm.62046	BG078369 Mm.62046 Signal Transduction
1500	transient 1 H3027C06 protein (C	receptor potential-related	Mus musculus transient receptor potential-related protein (ChaK), mRNA BG065092 Mm.33819 Energy/Metabolism	BG065092	Mm.33819	Energy/Metabolism
1502 1503	H3027E05 uridine pl	nosphorylase (Upp),	Mus musculus uridine phosphorylase (Upp), mRNA	BG065114 Mm.4610	1	Energy/Metabolism
1504 1505	H3027E07	alpha-enolase (2-phospho-D-glycerate H3027E07 hydrolase) (EC 4.2.1.11)	Mouse mRNA for alpha-enolase (2- phospho-D-glycerate hydrolase) (EC 4.2.1.11)	BG078408	Mm.90587	Mm.90587 Energy/Metabolism
1506 1507	H3027E08	•	Mouse mRNA for alpha-enolase (2- phospho-D-glycerate hydrolase) (EC 4.2.1.11)	BG078409	Мт.90587	BG078409 Mm.90587 Energy/Metabolism
1508 1509	H3027E09	o-D-glycerate	Mouse mRNA for alpha-enolase (2- phospho-D-glycerate hydrolase) (EC 4.2.1.11)	BG078410	Mm.90587	Mm.90587 Energy/Metabolism
1510 1511	H3027F02		Homo sapiens aspartyl-tRNA synthetase (DARS), mRNA		Mm.28693	BG078414 Mm.28693 Energy/Metabolism
1512 1513	H3027F07	"Murine MAP kinase kinase 6c mRNA, H3027F07 complete cds"		BG065128	Mm.14487	Signal Transduction
1514 1515	H3027F12	"Mus musculus Cctd gene for chaperonin containing TCP-1 delta H3027F12 subunit, complete cds"		BG078424 Mm.6821		Heat Shock/Stress

	Mm.38470 Heat Shock/Stress	Energy/Metabolism	Energy/Metabolism	BG065166 Mm.21908 Energy/Metabolism	Energy/Metabolism	Mm.19352 Signal Transduction	Apoptosis	Signal Transduction	Signal Transduction	Energy/Metabolism	BG078499 Mm.46396 Signal Transduction	Signal Transduction
	Mm.38470	Mm.5246	Mm.5246	Mm.21908	Mm.658	Mm.19352	BG078504 Mm.29028 Apoptosis	Mm.1034		Mm.15259	Mm.46396	Mm.2277
,	BG078439	BG078450	BG078451			BG065190	BG078504	BG078507	BG078497 Mm.930	BG065221	BG078499	BG065250
		Mus musculus peptidylprolyl isomerase A (Ppia), mRNA	Mus musculus peptidylprolyl isomerase A (Ppia), mRNA	Mus musculus BTB and CNC homology 2 (Bach2), mRNA	Mus musculus adenine nucleotide translocase-2 (Ant2) mRNA, complete cds		Mus musculus mRNA for death associated protein 3 (DAP-3 gene)			Mouse kidney ornithine decarboxylase mRNA, clone pODC16, 3 end		
	"Homo sapiens heat shock protein 75 (TRAP1), mRNA"	prolyl isomerase A (Ppia),	•	d CNC homology 2 (Bach2),	mrcleotide franslocase-2 (Ant2)	nusculus adenosine kinase (Adk),	Mus musculus mRNA for death H3008F01 death associated protein 3 (DAP-3 gene) associated protein 3 (DAP-3 gene)	"Mus musculus protein phosphatase 2 (formerly 2A), regulatory subunit A (PR H3028E04 65), alpha isoform (Ppp2r1a), mRNA"	"Mus musculus cathepsin L (Ctsl), mRNA"	ornithine decarboxylase, clone 6, 3 end	"Mus musculus RAB17, member RAS H3028F06 oncogene family (Rab17), mRNA"	"Mus musculus cathepsin H (Ctsh), mRNA"
	"Homo H3027H04(TRAP1	H3028A03	H3028A04 peptidyl	H3028A09BTB an	U3028B03 adenine	"Mus m	H3028F01	H3028E04	"Mus n H3028F04 mRNA	kidney H3028F05 pODCJ	H3028F06	"Mus n H3028H11 mRNA'
	1516	1518 1519	1520	1522 1523	1524	1526	1528	1530	1532	1534 1535	1536 1537	1538 1539

1540	peroxisom	al trans 2-enoyl CoA	Mus musculus peroxisomal trans 2- enoyl CoA reductase mRNA, complete cds	BG065254 Mm.29988	/m.29988	Energy/Metabolism
1541 1542 1543	H3029A05	optosis susceptibility protein	Mus musculus cellular apoptosis susceptibility protein mRNA, complete cds	BG065256 Mm.22417 Apoptosis	Mm.22417	Apoptosis
1544			Mus musculus natural resistance associated macrophage protein-2 (Nramp2) mRNA, C-terminal exon alternative solice variant, complete eds	BG065264	Mm.1304	Energy/Metabolism
1545	H3029B01 terminal	sculus chaperonin subunit 3		BG078552	Mm.27804	Heat Shock/Stress
1548	H3029C0	oiquinone) 1, 14.5kD,	Homo sapiens NADH dehydrogenase (ubiquinone) 1, subcomplex unknown, 2 (14.5kD, B14.5b) (NDUFC2), mRNA	BG078565	Mm.1893	Energy/Metabolism
1550	H3029C06					Heat Shock/Stress
1552	H3029E0	"Mus musculus Son of sevenless homolog 1, (Drosophila) (Sos1), H3029E04 mRNA"		BG065298	Mm.6357	Signal Transduction
1554	H3029G1	nuclear mitotic apparatus protein 1	Homo sapiens nuclear mitotic apparatus protein 1 (NUMA1), mRNA		Mm.102520	
1555	H3030A0	H3030A03 Mouse cDNA for heat shock protein 65		BG078626	Mm.1777	Heat Shock/Stress
1557	H3030B07	77			Mm.1776	Heat Shock/Stress
1559 1560	H3030C0	"Mus musculus activin receptor IIB H3030C05(Acvr2b), mRNA"	1	BG065366 Mm.8940	Mm.8940	Signal Transduction

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Signal Transduction	Energy/Metabolism	Energy/Metabolism	Energy/Metabolism		Mm.27953 Energy/Metabolism	BG078670 Mm.27953 Energy/Metabolism	Signal Transduction	Matrix/Structural Proteins	Energy/Metabolism	Energy/Metabolism	
Mm.1260		Mm.2635	Mm.2635	Mm.26022	Mm.27953	Mm.27953	Mm.22753	Matrix/S AW537455 Mm.142697 Proteins	Mm.43671	Mm.4069	20121
BG078650 Mm.1260		BG078663 Mm.2635	BG078664		BG078669	BG078670	BG078674	AW537455	BG078685	BG078689	00055000
		Mus musculus mRNA for pyruvate kinase M, complete cds	Mus musculus mRNA for pyruvate kinase M, complete cds		Human glycine decarboxylase (P- protein) mRNA	Human glycine decarboxylase (P- protein) mRNA		·	M.musculus partial cochlear mRNA (clone 20F5)	Mouse ATP synthase alpha subunit, complete cds	
"Mus musculus nucleoside diphosphate kinase A long form mRNA, complete cds"	H3030C11	H3030D10 pyruvate kinase M	H3030D11 pyruvate kinase M	Mm.26022	glycine decarboxylase (P-protein)	glycine decarboxylase (P-protein)	"Mus musculus cathepsin B (Ctsb), mRNA"	Homo sapiens protein phosphatase 3 (formerly 2B), catalytic subunit, beta isoform (calcineurin A beta) (PPP3CB), mRNA	H3030G01 Cochlear (clone 20F5)	ıthase alpha subunit	"Mus musculus Ca2+-independent phospholipase A2 long form mRNA,
"Mu kina H3030C06 cds"	H3030C11 H30300	H3030D10	H3030D11	H3030E04 Mm.260	H3030E05 glycine	H3030E06 glycine	"Mus m H3030E10 mRNA"	Homo s (former isoform H3030F09 mRNA	H3030G01	H3030G08 ATP syr	H3030G00
1561 1562	1563 1564	1565 1566	1567 1568	1569 1570	1571 1572	1573 1574	1575 1576	1577 1578	1579 1580	1581 1582	1583

					natin					
Signal Transduction	Signal Transduction	Mm.99776 Energy/Metabolism	BG065447 Mm.28219 Energy/Metabolism	Signal Transduction	Transcription/Chromatin	Energy/Metabolism	Energy/Metabolism	Energy/Metabolism	Mm.30158 Energy/Metabolism	Energy/Metabolism
Мт.8903	Mm.70272	Mm.99776	Mm.28219	Mm.27063		Mm.16763	Mm.2573	i	Mm.30158	Mm.972
BG078694	BG078710		BG065447	BG078729	BG078732 Mm.4063	BG065457	BG065475	BG065476 Mm.5289	BG078755	BG065478 Mm.972
Mus musculus branched chain ketoacid dehydrogenase kinase (Bckdk), mRNA	·	Homo sapiens cytosolic aminopeptidase P mRNA, complete cds BG078715	Rattus norvegicus PIS1 gene for phosphatidylinositol synthase, complete cds		·	Mus musculus aldolase 1, A isoform (Aldo1), mRNA	Mus musculus S-adenosylhomocysteine hydrolase (Ahcy), mRNA	Mus musculus glyceraldehyde-3- phosphate dehydrogenase (Gapd), mRNA	Homo sapiens fatty acid desaturase 1 (FADS1), mRNA	Homo sapiens ubiquinol-cytochrome c reductase core protein I (UQCRC1), mRNA
	"Mus musculus protein kinase C substrate 80K-H (Prkcsh), mRNA"	aminopeptidase P	Ratt phosphatidylinositol synthase cds	"Mus musculus putative intracellular signaling protein (Trip6) mRNA,			S-adenosylhomocysteine hydrolase	glyceraldehyde-3-phosphate	H3031E12 fatty acid desaturase 1 (FADS1),	ubiquinol-cytochrome c reductase core
H3030H02 Bckdk	"Mus mu H3031A11 substrate	H3031B06 cytosolic	H3031C05	· "Mus mu signaling	Mus mus	H3041D03 aldolase	S-adeno H3031F10 (Ahev)	H3031E11	H3031E12	ubiquino H3031F01 mrotein I
1585 1586	1587 1588	1589	1591	1593	1595	1597	1599	1601	1603	1605

1607		"Mus musculus heat shock protein 40 (HSPF1), mRNA"		BG065483	Mm.2982	Heat Shock/Stress
	H3031F12	ılus aspartate aminotransferase n 10 and 3-flank		BG078765	Mm.149089	BG078765 Mm.149089 Energy/Metabolism
	H3031G06	"Mus musculus heat shock protein, 110 H3031G06kDa (Hsp110), mRNA"		BG065493 Mm.1032		Heat Shock/Stress
1613 1614	H3031G08 Mm.932	Мт.93266			Mm.93266	Mm.93266 Energy/Metabolism
1615 1616	H3031G11	H+ transporting, lysosomal r proton pump) 9kD (ATP6H),	Homo sapiens ATPase, H+ transporting, lysosomal (vacuolar proton pump) 9kD (ATP6H), mRNA		Mm.22602	BG078776 Mm.22602 Energy/Metabolism
1617	H3031H09 5-3 exori		Mus musculus 5-3 exoribonuclease 2 (Xrn2), mRNA	BG065507 Mm.3065		Energy/Metabolism
1619 1620	H3032A01	ical protein PRO1197 77),	Homo sapiens hypothetical protein PRO1197 (PRO1197), mRNA	BG065511	Mm.24565	Mm.24565 Energy/Metabolism
1621	"Mus m 5 (gluc H3032A08(Hspa5)	"Mus musculus heat shock 70kD protein 5 (glucose-regulated protein, 78kD) (Hspa5), mRNA"		BG078795 Mm.918	Mm.918	Heat Shock/Stress
1623 1624	peroxis H3032A09PMP34	mal integral membrane protein	Mus musculus mRNA for peroxisonal integral membrane protein PMP34		Mm.306	Matrix/Structural Proteins
1625 1626	H3032A12	glucose-6-phosphate dehydrogenase X-13032A12 linked (G6pdx),	Mus musculus glucose-6-phosphate dehydrogenase X-linked (G6pdx), mRNA	BG078799	Mm.27210	BG078799 Mm.27210 Energy/Metabolism
1627 1628	H3032C01	H3032C01 cytochrome c gene (MC1)	Mouse cytochrome c gene (MC1)	BG078810	Mm.35389	BG078810 Mm.35389 Energy/Metabolism

1629	H3032C10	3-hydroxy-3-methylglutaryl-Coenzyme H3032C10 A reductase (HMGCR),	Homo sapiens 3-hydroxy-3- methylglutaryl-Coenzyme A reductase (HMGCR), mRNA	BG078816 Mm.2226	1 1	Energy/Metabolism
1631 1632	phosph H3032H02 (Pctpl),	atidylcholine transfer protein-like	dylcholine transfer protein-like Mus musculus phosphatidylcholine transfer protein-like (Pctpl), mRNA	BG078855	Mm.28896	Energy/Metabolism
1633 1634	H3033A10	R.norvegicus gene 6-phosphofructo-2-kinase/fructose-2,6-biphosphatase (EC H3033A102.7.1.105/EC 3.1.3.46)	R.norvegicus gene for 6-phosphofructo- 2-kinase/fructose-2,6-biphosphatase (EC 2.7.1.105/EC 3.1.3,46)	BG078872		Energy/Metabolism
1635	cytochr H3033B08(Cox4),	subunit IV	Mus musculus cytochrome c oxidase, subunit IV (Cox4), mRNA	BG078879	Mm.2136	Energy/Metabolism
1637 1638	H3033B11 serine pro	otease inhibitor 3 (Spi3),	Mus musculus serine protease inhibitor 3 (Spi3), mRNA	BG078882	Mm.147649	Mm.147649 Energy/Metabolism
1639	H3033C02	"Mus musculus serine/threonine kinase H3033C02(sak-a) mRNA, complete cds"		BG078885	Mm.3794	Signal Transduction
1641	ectonucle pyrophos H3033E10 (Enpp1),	ectonucleotide pyrophosphatase/phosphodiesterase 1 (Enpp1),	Mus musculus ectonucleotide pyrophosphatase/phosphodiesterase 1 (Enpp1), mRNA	BG065640	Mm.27254	Mm.27254 Energy/Metabolism
1643	ATP-bin H3034B11 (MDR/T.	ATP-binding cassette, sub-family B (MDR/TAP), member 12 ABC-me			Mm.143731	Mm.143731 Energy/Metabolism
1645	tumor nec H3035A01 protein 2	tumor necrosis factor, alpha-induced protein 2 (Tnfaip2),	Mus musculus tumor necrosis factor, apha-induced protein 2 (Tufaip2), mRNA	BG065761	Mm.4348	Apoptosis
1647	H3035B03	H3035B03 Mm.32746			Mm.32746	Energy/Metabolism
1649 1650	Mouse g H3035D08 receptor	Mouse gene for beta-2-adrenergic receptor		BG079067	Mm.5598	Signal Transduction

Signal Transduction	Signal Transduction	Signal Transduction		BG079254 Mm.104955 Energy/Metabolism	Mm.154704 Signal Transduction	Heat Shock/Stress	Mm.113278 Energy/Metabolism	BG079313 Mm.27703 Energy/Metabolism
Мт.2411	Mm.3181	Mm.4772	Mm.27828	Mm.104955	Mm.154704	Мт.21761	Mm.113278	Mm.27703
BG065817	BG079136	BG079186		BG079254	BG066005	BG079287	BG079311	BG079313
			Homo sapiens kinesin-like 6 (mitotic centromere-associated kinesin) (KNSL6), mRNA	Mus musculus serine protease inhibitor, Kunitz type 1 (Spint1), mRNA			Homo sapiens potassium voltage-gated channel, delayed-rectifier, subfamily S, member 3 (KCNS3), mRNA	Homo sapiens phosphoribosyl pyrophosphate synthetase-associated protein 2 (PRPSAP2), mRNA
"Mus musculus RNA-binding protein isoform G3BP-2a (G3BP2) mRNA, H3035F02 [complete cds"	"Mus musculus Rho guanine nucleotide exchange factor (GEF) 1 (Arhgef1), mRNA"	"Mus musculus Iysophosphatidic acid receptor (vzg-1) mRNA, complete cds"	kinesin-like 6 (mitotic centromere- H3037B01 associated kinesin) (KNSL6),	serine protease inhibitor, Kunitz type 1 (Spint1),	"Mus musculus cAMP-specific phosphodiesterase 4A (Pde4a) gene, exons 2 through 8 and PDE4A isoform 1 exon 1"	"Mus musculus interferon alpha responsive protein (15 kDa) (Iffg15), mRNA"	potassium voltage-gated channel, delayed-rectifier, subfamily S, member H3038C07[3 (KCNS3),	phosphoribosyl pyrophosphate synthetase-associated protein 2 H3038C09((PRPSAP2),
H3035F02	"Mus m exchang H3036B11 mRNA"	H3036H01	H3037B01	serine p H3037F06 (Spint1	"Mus n phosph exons 2 H3037G121 exon	"Mus m respons H3038A06 mRNA'	H3038C07	H3038C09
1651	1653	1655	1657	1659 1660	1661 1662	1663 1664	1665 1666	1667 1668

1669	PRKC, application	ooptosis, WT1, regulator	Homo sapiens PRKC, apoptosis, WT1, regulator (PAWR), mRNA	BG079331	Mm.6617	Apoptosis
1671	Mus mu H3048F05	Mus musculus growth differentiation		BG079333	Mm.4213	Signal Transduction
1673 1674	H3038E10	ansferase COQ3 (COQ3),	Homo sapiens methyltransferase COQ3 (COQ3), mRNA	BG066071	Mm.5662	Energy/Metabolism
1675	"Mus mu H3039A05 signaling	"Mus musculus regulator of G-protein signaling 11 mRNA, partial cds"		BG079374	Mm.13264	Signal Transduction
1677	"Mus mu receptor H3039B09mRNA"	"Mus musculus serine/threonine kinase receptor associated protein (Strap), mRNA"		BG066121	Mm.22584	Signal Transduction
1679	H3039C11	"Mus musculus receptor tyrosine kinase (Dtk) mRNA, complete cds"		BG066134	Mm.2901	Signal Transduction
1681	H3039D02	(NARS),	Homo sapiens asparaginyl-tRNA synthetase (NARS), mRNA	BG079401	Mm.29192	Mm.29192 Energy/Metabolism
1683	H3039E07	isocitrate dehydrogenase 3 (NAD+) beta: (IDH3B),	dehydrogenase 3 (NAD+) beta Homo sapiens isocitrate dehydrogenase 3 (NAD+) beta (DH3B), mRNA	C78231	Mm.29590	Mm.29590 Energy/Metabolism
1685	H3039E08	"Mus musculus SH3-containing protein		BG079417 Mm.4165	Mm.4165	Signal Transduction
1687	H3039F05 ornithine	ornithine aminotransferase (Oat),	Mus musculus ornithine aminotransferase (Oat), mRNA	BG079424	Mm.42187	BG079424 Mm.42187 Energy/Metabolism
1689 1690	M.musc H3039G04(SERS)	M.musculus seryl-tRNA synthetase (SERS) , 5 end	M.musculus seryl-tRNA synthetase (SERS) mRNA, 5 end	BG079434	Mm.28688	BG079434 Mm.28688 Energy/Metabolism

1691 1692	H3039G12	H3039G125-3 exoribonuclease 1 (Xm1),	Mus musculus 5-3 exoribonuclease 1 (Xm1), mRNA	BG079441	Mm.5703	Energy/Metabolism
1693 1694	"Mus m H3039H01 mRNA'	"Mus musculus calponin 2 (Cnn2), mRNA"		BG079442	Mm.21776	Mm.21776 Signal Transduction
1695 1696	H3039H05	Mus musculus putative CCAAT binding factor 1 (mCBF) mRNA, alternatively H3039H05 spliced transcript mCBF1, complete cds		BG079446	Mm.24169	
1697 1698	adenine H3040A04 (APRT)	adenine phosphoribosyltransferase (APRT)	Mouse adenine phosphoribosyltransferase (APRT), complete cds	BG079455	Mm.1786	Energy/Metabolism
1699 1700	H3040E11	H3040E11 palmitoyl-protein thioesterase (Ppt),	Mus musculus palmitoyl-protein thioesterase (Ppt), mRNA		Mm.153740	
1701 1702	H3041A02	NADH dehydrogenase (ubiquinone) 1 alpha subcomplex, 10 (42kD) H3041A02(NDUFA10),	Homo sapiens NADH dehydrogenase (ubiquinone) 1 alpha subcomplex, 10 (42kD) (NDUFA10), mRNA	BG079485	Mm.28293	Enerov/Metaholism
1703 1704	H3041A04	H3041A04 lysyl-tRNA synthetase (KARS),	Homo sapiens lysyl-tRNA synthetase (KARS), mRNA		Mm.29949	Mm.29949 Energy/Metabolism
1705 1706	mevalon H3041B11 decarbox	ate pyrophosphate tylase	Rattus norvegicus mevalonate pyrophosphate decarboxylase mRNA, complete cds	BG079503	Mm.28146	Mm.28146 Energy/Metabolism
1707 1708	H3041F09 glucose	transporter (GLUTX1 gene)	Mus musculus mRNA for glucose transporter (GLUTX1 gene)	BG066327 Mm.7241	Mm.7241	Energy/Metabolism
1709 1710	solute ca H3041G03 transport	rrier family 15 (H+/peptide er), member 2 (Slc15a2),	Mus musculus solute carrier family 15 (H+/peptide transporter), member 2 (Slc15a2), mRNA	BG079545	Mm.63479	BG079545 Mm.63479 Energy/Metabolism

1711	profease H3041G08subunit.	(prosome, macropain) 26S ATPase 5 (Psmc5),	Mus musculus protease (prosome, macropain) 26S subunit, ATPase 5 (Psmc5), mRNA	BG079550	Mm.665	Energy/Metabolism
		stocompatibility complex	Mus musculus major histocompatibility			
1713		i, RPSZs, NADH 29, KIFCI, Fas- VG1, tapasin, BING4, beta 1,3-	sse, NG29, KIFCI, BINGI, tapasin, BING4, beta 1,3-			
1714	H3041H03	H3041H03 galactosyl transfera galactosyl transfera Homo sapiens BCL2-associate H3042 405 RCI 2-associated athanogene 4 (BAG4), athanogene 4 (BAG4), mRNA	D.	BG066354	Mm.27102 Apoptosis	Energy interacousing Apoptosis
1717	H3042C08	"Mus musculus dual specificity H3042C08 phosphatase 10 (Dusp10), mRNA"		BG079592	Mm.34912	Mm.34912 Signal Transduction
1719	H3042D02	milar ag3),	Mus musculus brain cDNA, clone MNCb-2243, similar to Mus musculus Bcl2-associated athanogene 3 (Bag3), mRNA	BG079597 Mm.28373 Apoptosis	Mm.28373	Apoptosis
1721	H3042F12		Mus musculus serine protease inhibitor 4 (Spi4), mRNA	BG079624	Mm.3093	Energy/Metabolism
1723 1724	H3042G07	H3042G07 Mouse heat-shock protein hsp84 mRNA		BG079631	Mm.2180	Heat Shock/Stress
1725 1726	"Mus mu 1, relate H3043A03 mRNA"	"Mus musculus casein kinase II, alpha 1, related sequence 4 (Csnk2a1-rs4), mRNA"		BG066436	Mm.23692	Mm.23692 Signal Transduction
1727 1728	Mus bind H3043F09 cds	Mus musculus GNB3 gene for GTP-binding protein beta3 subunit, complete cds		BG066499	Mm.22228	
1729 1730	H3043F12	H3043F12 ferrochelatase (Fech),	Mus musculus ferrochelatase (Fech), mRNA	BG066502	Mm.1070	

													
Mm.36431 Heat Shock/Stress	100 H2 H2 100 H2 100 H2 100 H2 100 H2 100 H2 100 H2 100 H2 100 H2 100 H2		Luci Bytweiabulisii	Mm 23710 Energy/Metabolism	Ticoponosial disease	Mun.155195)Energy/Metabolism	Energy/Metabolism	Mm.25054 Energy/Metabolism	Energy/Metabolism		Mm.29193 Apoptosis	Grown' Mart. 11	BG066807 Mm.28867 Energy/Metabolism
I .	1			Mm 23710	Mr. 16510	100 I I I I I I I I I I I I I I I I I I	Mm.7060	Mm.25054	Mm.21880	7775 mJV	Mm.29193	Mm 20847	Mm.28867
BG079699	אכטעעעניט	BG066673	BG070700	06161000	B/3070700	210000	C186/009		BG079822	BG170861		BG079880	3G066807
	Mus musculus protease (prosome, macropain) 28 subunit, alpha (Psme1), mRNA	Mus musculus A10 mRNA, partial cds BG066673	Homo sapiens cytosolic aminoperidase P mRNA complete eds BG070700		Mus musculus solute carrier family 12, member 7 (Slc12a7) mRNA	Homo sapiens mRNA for NAALADase	a process		Mus musculus serine protease OMI (Omi), mRNA	Homo sapiens DKFZP5660084 protein (DKFZP5660084), mRNA	l cell death		dase II
"Mus musculus Cctd gene for chaperonin containing TCP-1 delta H3044B01 subunit, complete cds"	protease (prosome, macropain) 28 H3044E06 subunit, alpha (Psme1),	H3044G06 A10 , partial cds	H3045B02 cytosolic aminopeptidase P	H3045B03 Mm.23710	solute carrier family 12, member 7 H3045B12 (S1c12a7),	ase II protein	77		H3045E05 serine protease OMI (Omi),	DKFZP5660084 protein H3046A03 (DKFZP5660084),	programmed cell death protein 7 H3046A12(Pdcd7),	ntenyl-diphosphate delta isomerase	H3047B07 tripeptidyl peptidase II (Tpp2),
H3044B	H3044E	H3044G	H3045B	H3045B	H3045B	H3045D(H3045D(H3045E0	H3046A0	H3046A1	isoper H3046C10 (IDI1)	H3047B0
1731 1732	1733 1734	1735 1736	1737 1738	1739 1740	1741 1742	1743	1745 1746	1747	1748	1749 1750	1751 1752	1753 1754	1755 1756

1757	חמראסבט	U2047D01 Ros faurus creatine kinase	Bos taurus mRNA for creatine kinase, complete cds	BG079988	Mm.970	Energy/Metabolism
1759	gene enco H3047D05 exons 5.6	hydratase,	te encoding encyl-CoA	BG079992	Mm.24452	Mm.24452 Energy/Metabolism
1761	vacuolar H3047F02 ATPase)	adenosine trisphosphatase (V-	(g)	BG079910	Mm.25079	Mm.25079 Energy/Metabolism
1763	H3047G12	tein (C7-1)	Rattus norvegicus C7-1 protein (C7-1) mRNA, complete cds	C80679	Mm.21961	Energy/Metabolism
1765	H3048A05	rboxyl CMT),	Homo sapiens isoprenylcysteine carboxyl methyltransferase (ICMT), mRNA	BG080028	Mm.44565	Energy/Metabolism
1767	cytochrol H3048B11 (Cox7c),	ounit VIIc	Mus musculus cytochrome c oxidase, subunit VIIc (Cox7c), mRNA	BG080036	Mm.24165	BG080036 Mm.24165 Energy/Metabolism
1769	small z H3048E06 (Ddo2)	inc finger-like protein DDP2	Mus musculus small zinc finger-like protein DDP2 (Ddp2) mRNA, complete cds	~ BG080049	Mm.30718	Mm.30718 Energy/Metabolism
1771	H3048G11	lin reductase B (flavin reductase PH)) (BLVRB),	Homo sapiens biliverdin reductase B (flavin reductase (NADPH)) (BLVRB), mRNA	BG066946	Mm.24021	Mm.24021 Energy/Metabolism
1773	"Mus mi H3049D07 mRNA"	"Mus musculus Janus kinase 2 (Jak2),		BG080085	Mm.25112	Mm.25112 Signal Transduction
1775	H3049F02 carbonyl	reductase 3 (CBR3),	Homo sapiens carbonyl reductase 3 (CBR3), mRNA	BG067014	Mm.4512	Energy/Metabolism
1777 1778	solute ca (monoca H3049G02 member	rrier family 16 rboxylic acid transporters), 1 (Slc16a1),	Mus musculus solute carrier family 16 (monocarboxylic acid transporters), member 1 (Slc16a1), mRNA	BG067025	Mm.9086	Energy/Metabolism
1779 1780	"Mus m leukemie H3049G04 _{mRNA} "	"Mus musculus B-cell leukemia/lymphoma 10 (Bcl10), mRNA"		BG067027	Mm.28782	BG067027 Mm.28782 Signal Transduction

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Mus musculus PGES mRNA for prostaglandin E synthase prostaglandin E synthase BG067030
Mus musculus small zinc finger-like
Homo sapiens putative putative dimethyladenosine transferase dimethyladenosine transferase (HSA9761), mRNA
Homo sapiens diaphorase (NADH) (cytochrome b-5 (cytochrome b-5 reductase) (DIA1), mRNA
hydroxysteroid 17-beta dehydrogenase 4 Mus musculus hydroxysteroid 17-beta dehydrogenase 4 (Hsd17b4), mRNA
solute carrier family 7 (cationic amino Mus musculus solute carrier family 7 acid transporter, y+ system), member 2 (cationic amino acid transporter, y+ system), member 2 (Slc7a2), mRNA system), member 2 (Slc7a2), mRNA
ssculus interferon regulatory
H3050H11 Mm.25374
H3051C06
"Mus musculus phospholipase D2 gene,
H3051C07 exons 13 through 25 and complete cds"

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	Mm.28344 Signal Transduction	Mm.29192 Energy/Metabolism	Signal Transduction	Mm.140599 Energy/Metabolism	Mm.25125 Energy/Metabolism	Signal Transduction	Signal Transduction	Signal Transduction	Energy/Metabolism	Mm.86738 Signal Transduction	
	Mm.28344	Мт.29192	Mm.28262	Mm.140599	Mm.25125		Mm.4356	Mm.29545	Mm.2124	Mm.86738	
	BG080373	BG080379	BG080390	BG067326	BG080402	BG080409	BG067357	BG067384	BG083930	BG080476	
		Homo sapiens asparaginyl-tRNA synthetase (NARS), mRNA		Mus musculus tyrosine hydroxylase (Th), mRNA	Homo sapiens phosphoribosyl pyrophosphate synthetase-associated protein 1 (PRPSAP1), mRNA			•	Mus musculus citrin (Slc25a13) mRNA, complete cds		
	"Homo sapiens rab3 GTPasc-activating protein, non-catalytic subunit (150kD) (RAB3-GAP150), mRNA"		"Mus musculus regulator of 'G-protein H3053A12/signaling 2 mRNA, complete cds"	tyrosine hydroxylase (Th),	phosphoribosyl pyrophosphate synthetase-associated protein 1 H3053C02(PRPSAP1),	"Mus musculus nicotinic acetylcholine receptor alpha 5 subunit (Acra5)	"Mus musculus calponin 1 (Cnn1), mRNA"	"Mus musculus polynucleotide kinase H3053G123'- phosphatase (Pnkp), mRNA"	H3053H06 citrin (Slc25a13)	"Mus musculus folate receptor 3	
	"Homo protein, H3052H05(RAB3-	H3052H11	H3053A12	H3053B07 tyrosine	H3053C02	"Mus m receptor H3053C09mRNA,	"Mus m H3053E04 mRNA"	H3053G12	H3053H06	H3054B05	
	1823	1825	1827	1829 1830	1831	1833	1835	1837 1838	1839 1840	1841	

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"Homo sapiens rab3 GTPase-activating protein, non-catalytic subunit (150kD) H3052H05(RAB3-GAP150), mRNA"	"Homo sapiens rab3 GTPase-activating protein, non-catalytic subunit (150kD) (RAB3-GAP150), mRNA"			BG080373	Mm.28344	Mm.28344 Signal Transduction
nyl-tRNA synthetase (NARS),	nyl-tRNA synthetase (NARS),		Homo sapiens asparaginyl-tRNA synthetase (NARS), mRNA	BG080379	Мт.29192	Energy/Metabolism
"Mus musculus regulator of G-protein H3053A12 signaling 2 mRNA, complete cds"	"Mus musculus regulator of G-protein isignaling 2 mRNA, complete cds"			BG080390	Мт.28262	Signal Transduction
H3053B07 tyrosine hydroxylase (Th),	lyrosine hydroxylase (Th),	Т	Mus musculus tyrosine hydroxylase (Th), mRNA	BG067326	Mm.140599	Mm.140599 Energy/Metabolism
phosphoribosyl pyrophosphate synthetase-associated protein 1 H3053C02 (PRPSAP1),	phosphoribosyl pyrophosphate synthetase-associated protein 1 (PRPSAP1),		Homo sapiens phosphoribosyl pyrophosphate synthetase-associated protein 1 (PRPSAP1), mRNA	BG080402	Mm.25125	Mm.25125 Energy/Metabolism
"Mus musculus nicotinic acetylcholine receptor alpha 5 subunit (Acra5) H3053C09 mRNA, partial cds"	"Mus musculus nicotinic acetylcholine receptor alpha 5 subunit (Acra5)			BG080409		Signal Transduction
"Mus musculus calponin 1 (Cnn1), H3053E04 mRNA"	"Mus musculus calponin 1 (Cnn1),			BG067357	Mm.4356	Signal Transduction
"Mus musculus polynucleotide kinase H3053G12[3- phosphatase (Pnkp), mRNA"	"Mus musculus polynucleotide kinase 23'- phosphatase (Pnkp), mRNA"	l i	·	BG067384	Mm.29545	Signal Transduction
H3053H06 citrin (Slc25a13)	Soitrin (Slc25a13)		Mus musculus citrin (Slc25a13) mRNA, complete cds	BG083930	Mm.2124	Energy/Metabolism
"Mus musculus folate receptor 3 H3054B05 (Folbp3) mRNA, complete cds"	"Mus musculus folate receptor 3 5(Folbp3) mRNA, complete eds"			BG080476	Mm.86738	Signal Transduction
H3054B12\Nrf2			Mus musculus p45 NF-E2 related factor 2 (Nrf 2) mRNA, complete cds	BG067417	Mm.1025	

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·	Energy/Metabolism	Signal Transduction	Signal Transduction	Mm.21583 Energy/Metabolism	Mm.87581 Energy/Metabolism	Apoptosis	Mm.90076 Energy/Metabolism	Energy/Metabolism
	Mm.28269	Mm.2038		Mm.21583	Mm.87581	Mm.1803	Mm.90076	
		BG080577	BG080578	BG080583	BG080587	BG080604	BG080616	BG080623 Mm.684
Mus musculus major histocompatibility locus class III region:butyrophilin-like protein gene, partial cds, Notch4, PBX2, RAGE, lysophatidic acid acyl transferase-alpha, palmitoyl-protein				Homo sapiens ATP/GTP-binding protein (HEAB), mRNA	Homo sapiens aldolase B, fructose- bisphosphate (ALDOB), mRNA	Mus musculus TRAF family member associated NF-kappa B activator (TANK) mRNA, complete cds	Mus musculus delta-aminolevulinate dehydratase (Lv), mRNA	Mus musculus dipeptidyl peptidase I precursor mRNA, complete cds
us class III ein gene, tAGE, erase-alpha,	inhibitor of kappa light polypeptide gene enhancer in B-cells, kinase beta	"Mus musculus Ras-GTPase-activating protein SH3-domain binding protein H3055C04 (G3bp-pending), mRNA"	"Homo sapiens inositol 1,3,4 triphosphate 5/6 kinase (ITPKI), mRNA"	H3055C10 ATP/GTP-binding protein (HEAB),	aldolase B, fructose-bisphosphate (ALDOB),	TRAF family member associated NF- leanna B activator (TANK)		H3055G02 dipeptidyl peptidase I precursor
	H3033B08	H3055C04	"Homo s triphospl H3055C05 mRNA"	H3055C10	aldolase B H3055D02(ALDOB)	TRAF H3055F07 kanna	H3055F07	H3055G02
1863	1865	1867	1869	1871	1873	1875	1877	1879 1880

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1881 1882	H3056A09	"M.musculus mRNA for inositol 1,4,5-H3056A09 trisphosphate receptor (type 2)"		BG067589	Mm.7800	Signal Transduction
	H3056C11		Homo sapiens DKFZP564O2082 protein (DKFZP564O2082), mRNA		Mm.21826	
	"Homo s H3056D03 (CNN3),	"Homo sapiens calponin 3, acidic (CNN3), mRNA"	•	BG080757 Mm.22171		Signal Transduction
	H3056G10	cin 2 (TRX2)	Homo sapiens thioredoxin 2 (TRX2) mRNA, complete cds	BG080633	Mm.3533	Energy/Metabolism
	H3056H05	Н3056Н05 Н3056Н05				Energy/Metabolism
1891 1892	"Mus mu devélopn H3057C01 mRNA"	"Mus musculus interferon-related developmental regulator 1 (ffrd1), mRNA"		BG067699	Mm.168	Heat Shock/Stress
1893	H3057C08 GPX3	GPX3	•		Mm.7156	Energy/Metabolism
1895 1896	H3057C10 bisphosp	hate 3-nucleotidase 1 (Bpnt1),	Mus musculus bisphosphate 3- nucleotidase 1 (Bpnt1), mRNA	BG067708	Мт.18096	Mm.18096 Energy/Metabolism
1897 1898	acetyl-Cc H3057D09(ACACB	oenzyme A carboxylase beta	Homo sapiens acetyl-Coenzyme A carboxylase beta (ACACB), mRNA	BG067718	Mm.88548	Mm.88548 Energy/Metabolism
1899 1900	"Mus mu phosphat H3057E05 (Ptpn21),	"Mus musculus protein tyrosine phosphatase, non-receptor type 21 (Ptpn21), mRNA"		BG080698	Mm.4420	Signal Transduction
1901	H3057F01	"Mus musculus casein kinase 1, epsilon (Csnk1e), mRNA"		BG080793	Мт.30199	Mm.30199 Signal Transduction
1903 1904	"Mus m H3057H04mRNA,			BG080820	Mm.46746	BG080820 Mm.46746 Heat Shock/Stress

Energy/Metabolism	Apoptosis	Apoptosis	Signal Transduction	Mm.27307 Signal Transduction	Mm.143646 Energy/Metabolism	Mm.34957 Energy/Metabolism	BG067869 Mm.31378 Signal Transduction	Mm.142839 Signal Transduction	Signal Transduction	Mm.24174 Energy/Metabolism
Mm.5045	BG080836 Mm.29026 Apoptosis	Mm.25988 Apoptosis	Mm.482	Mm.27307	Mm.143646	Mm.34957	Мт.31378	Mm.142839	BG067874 Mm.28598	Mm.24174
C87415		BG080862	BG080846	BG080868	BG080898	BG067865	BG067869	BG080928	BG067874	BG080951
Mus musculus solute carrier family 16 (monocarboxylic acid transporters), member 2 (Slc16a2), mRNA	Mus musculus Fas-interacting Fas-interacting serine/threonine kinase 3 (Fist3) mRNA, (Fist3)	Mus musculus Bcl2-like 10 (Bcl2110), mRNA			Mus musculus transient receptor potential-related protein (ChaK), mRNA BG080898	Mus musculus sodium bicarbonate bicarbonate cotransporter isom 3 (kNBC-3), mRNA		,		Homo sapiens alanyl-tRNA synthetase (AARS), mRNA
solute carrier family 16 (monocarboxylic acid transporters), H3057H06 member 2 (Slc16a2),	Fas-interacting scrine/threonine kinase 3 (Fist3)	H3058C03 Bcl2-like 10 (Bcl2110),	"Mus musculus Jun oncogene (Jun), mRNA"	"Mus musculus guanine nucleotide binding protein (G protein), gamma 3 subunit (Gng3), mRNA"		sodium bicarbonate cotransporter isom 3 (kNBC-3),	"Homo sapiens regulator of G-protein H3059B02 signalling 12 (RGS12), mRNA"	"Mus musculus protein kinase C delta mRNA, complete cds"	"Mus sp. JAK1 protein tyrosine kinase mRNA, complete cds"	H3059D07 alanyl-tRNA synthetase (AARS),
H3057H06	Fas-int H3058A10(Fist3)	H3058C03	"Mus II H3058C09 mRNA'	"Mus m binding H3058D09 subunit	transier H3058G06 protein	sodium H3059A10 (kNBC	H3059B02	"Mus m H3059B03 mRNA,	"Mus sr H3059B07 mRNA,	H3059D07
1905 1906	1907	1909 1910	1911 1912	1913 1914	1915 1916	191 <i>7</i> 1918	1919 1920	1921 1922	1923 1924	1925 1926

1927	H3059D11	H3059D11KIAA1093 protein, partial cds	Homo sapiens mRNA for KIAA1093 protein, partial cds	BG067897	Mm.22829	Apoptosis
	H3050F12	rotein		BG067921	Mm.15793	Signal Transduction
	H3059G11 intronless	gene	Mouse intronless glutamine synthetase gene, complete cds	BG067932	Mm.41660	Energy/Metabolism
1933	H3060A10		M.musculus mRNA for arachidonate epidermis-type 12(S)-lipoxygenase	BG067951	Mm.1122	Energy/Metabolism
1935	H3060B09	"Homo sapiens mitogen-activated protein kinase kinase kinase kinase 5 (MAP4K5), mRNA"		BG081019	Mm.24022	Signal Transduction
1937	H3060D04	H3060D04Mm.24594			Mm.24594	Energy/Metabolism
1939	H3060H04	"Mus musculus JNK-binding protein 1 H3060H04(Inkbp1-pending), mRNA"		BG081077	Mm.25540	Mm.25540 Signal Transduction
1941	Н3060Н07	H3060H07\Mm.25580			Mm.25580	Mm.25580 Energy/Metabolism
1943 1944	UDP-N-galactoss galactoss acetylgal H3061B06(Galnt3),	UDP-N-acetyl-alpha-D-galactosamine:polypeptide N-acetylgalactosaminyltransferase 3 (Galnt3),	Mus musculus UDP-N-acetyl-alpha-D-galactosamine:polypeptide N-acetylgalactosaminyltransferase 3 (Galnt3), mRNA.	BG068045	Mm.38441	Mm.38441 Energy/Metabolism
1945	H3061E04 cvsteine	cysteine dioxygenase, type I (CDO1),	steine dioxygenase, nRNA	BG081135	Mm.29996	Mm.29996 Energy/Metabolism
1947 1948	H3061H08 guanine	guanine deaminase (Gda),	Mus musculus guanine deaminase (Gda), mRNA	BG081171		Energy/Metabolism
1949 1950	H3062C04	H3062C04 spermine synthase (Sms),	Mus musculus spermine synthase (Sms), mRNA	BG081202		Energy/Metabolism

1951 1952	H3062E02	H3062E02 isocitrate dehydrogenase	Mus musculus isocitrate dehydrogenase mRNA, complete cds	BG081213	Mm.18213	Mm.18213 Energy/Metabolism
1953 1954	H3062H07	"Mus musculus inositol 1,4,5- H3062H07 triphosphate receptor 1 (Itpr1), mRNA"		BG081243	Mm.2726	Signal Transduction
1955 1956	H3062H08	H3062H08Mm.11827			Mm.11827	Mm.11827 Energy/Metabolism
1957 1958	"Mus m H3063A03 mRNA"	"Mus musculus calpain 7 (Capn7), mRNA"		BG081250	Мт.142370	Mm.142370 Signal Transduction
1959 1960	H3063A08 Prsc1	Prsc1	Mus musculus protease, cysteine, 1 (Prsc1), mRNA	BG068219	Mm.17185	Mm.17185 Energy/Metabolism
1961 1962	H3063A09	"Mus musculus novel ras effector 1 H3063A09 (Nore1-pending), mRNA"		BG068220	Mm.10133	Signal Transduction
1963 1964	H3063C09 beta-1,4	galactosyltransferase VI	Mus musculus beta-1,4- galactosyltransferase VI mRNA, complete cds	BG081279	Mm.26364	BG081279 Mm.26364 Energy/Metabolism
1965 1966	H3063H10	H3063H10LAT2 (Slc7a8)	Mus musculus mRNA for LAT2 protein (Slc7a8 gene)	BG068299	Mm.27830	Mm.27830 Energy/Metabolism
1967 1968	"Mus m H3064C02 1 (Lats)	"Mus musculus large tumor suppressor 1 (Lats1) mRNA, partial cds"		BG081357	Mm.35642	BG081357 Mm.35642 Signal Transduction
1969 1970	kinesin- H3064C04 protein	like 5 (mitotic kinesin-like l) (KNSL5),	Homo sapiens kinesin-like 5 (mitotic kinesin-like protein 1) (KNSL5), mRNA		Mm.28386 Cell Cycle	Cell Cycle
1971 1972	vacuola H3064E02 subunit	r adenosine triphosphatase A gene	Mus musculus vacuolar adenosine triphosphatase subunit A gene, complete cds	3G081377	Mm.29771	BG081377 Mm.29771 Energy/Metabolism
1973 1974	H3064H04	H3064H04 HMG box protein	Mus musculus HMG box protein mRNA, complete cds		Mm.41766	Mm.41766 Transcription/Chromatin

BG081448 Mm.24381 Signal Transduction	BG081451 Signal Transduction	BG068432 Mm.144498 Signal Transduction	Mus musculus UDP-glucose dehydrogenase (Ugdh), mRNA BG068439 Mm.10709 Energy/Metabolism	Mus musculus heme oxygenase-2 BG081591 Energy/Metabolism	Mus musculus potassium channel modulatory factor DEBT-91 (Debt91), BG081603 Mm.29194 Energy/Metabolism	BG081616 Mm.3058 Signal Transduction	BG068616 Mm.29014 Signal Transduction	Homo sapiens kinesin-like 5 (mitotic kinesin-like protein 1) (KNSL.5), mRNA) Mm.28386 Cell Cycle	A C
"Mus musculus calmodulin dependent phosphatase catalytic subunit (Cam-Prp) H3065C08ImRNA, 3' end"	"Mus musculus Chetk-alpha gene for choline/ethanolamine kinase-alpha,	"Homo sapiens RAP1A, member of RAS oncogene family (RAP1A), H3065D10 mRNA"	cose dehydrogenase (Ugdh),		n channel modulatory factor 1 (Debt91),	"Mus musculus phosphoinositide 3-kinase regulatory subunit p85alpha	"Mus musculus RAS-related C3 botulinum substrate 1, guanine nucleotide exchange factor 1 (Raegefl- H3067E02 pending), mRNA"	otic kinesin-like),	
1975 H. 1976									

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1995	H3068F03	ornithine decarboxylase antizyme H3068F03 inhibitor (Oazi),	Mus musculus ornithine decarboxylase antizyme inhibitor (Oazi), mRNA	BG081746	Mm.104010	BG081746 Mm.104010 Energy/Metabolism
1997 1998	H3069C09	"Mus musculus Rho-associated colled- H3069C09[coil forming kinase 1 (Rock1), mRNA"	·	BG081800	Mm.6710	Signal Transduction
1999	H3069D09		Homo sapiens cDNA FLJ12814 fis, clone NT2RP2002520, weakly similar to Homo sapiens transcription factor RFX-B (RFXB) mRNA	BG081823	Mm.27228	Mm.27228 Energy/Metabolism
2001 2002	choline H3069G01 (CEPT)	ethanolaminephosphotransferase),	Homo sapiens choline/ethanolaminephosphotransferase (CEPT1), mRNA	BG081849	Mm.14816	Mm.14816 Energy/Metabolism
2003 2004	H3070A09	RHOA proto-oncogene multi-drug- H3070A09 resistance protein , 3 end	Homo sapiens RHOA proto-oncogene multi-drug-resistance protein mRNA, 3 end	BG081880 Mm.757	Mm.757	Energy/Metabolism
2005 2006	quinoid H3070A12 (QDPR)	dihydropteridine reductase	Homo sapiens quinoid dihydropteridine reductase (QDPR), mRNA	BG081883	Мт.30204	Mm.30204 Energy/Metabolism
2007 2008	partial 1 H3070B09 protein,	Kcnq1 gene potassium channel exons 10-14	Mus musculus partial Kenq1 gene for potassium channel protein, exons 10-14 BG081892	BG081892		Energy/Metabolism
2009 2010	H3070C08	hydroxysteroid dehydrogenase-1, H3070C08 delta<5>-3-beta (Hsd3b1),	Mus musculus hydroxysteroid dehydrogenase-1, delta<5>-3-beta (Hsd3b1), mRNA	BG081903	Mm.16941	Energy/Metabolism
2011 2012	H3070E04	ATP-specific succinyl-CoA synthetase (H3070E04 beta subunit (Scs), partial cds	Mus musculus ATP-specific succinyl- CoA synthetase beta subunit (Scs) mRNA, partial cds	BG068897	Mm.19154	Mm.19154 Energy/Metabolism
2013 2014	H3071A03 ATPase,	class VI, type 11A (Atp11a),	Mus musculus ATPase, class VI, type 11A (Atp11a), mRNA	BG081967	Mm.148756	Mm.148756Energy/Metabolism

2015 2016	"Homo s RAS on H3071G11 mRNA"	"Homo sapiens RAP2B, member of RAS oncogene family (RAP2B), mRNA"		BG082041	Mm.26939	BG082041 Mm.26939 Signal Transduction
2017	H3071H04	3-phosphate acyltransferase	Mouse glycerol-3-phosphate acyltransferase mRNA, complete cds		Mm.87773	
2018 2019	H3072A08		Mus musculus glyceraldehyde-3- phosphate dehydrogenase (Gapd), mRNA	BG082061		Energy/Metabolism
2020 2021	H3072B11	"Mus musculus PLC-L2 mRNA for H3072B11 phospholipase C-L2, complete cds"		BG069051	Mm.28034	Mm.28034 Signal Transduction
2022 2023	H3072F03	H3072F03 Mm.22651			Mm.22651	Energy/Metabolism
2024 2025	potassiur subfamil H3072G09 (Kenh1),	n voltage-gated channel, y H (eag-related), member 1	Mus musculus potassium voltage-gated channel, subfamily H (eag-related), member I (Kcnhl), mRNA	BG069106	Mm.4489	Energy/Metabolism
2026 2027	H3073F09	H3073F09 Mm.27123			Mm.27123	Mm.27123 Energy/Metabolism
2028 2029	H3073F10	"Mus musculus 80kDa m-calpain H3073F10 subunit (calp80) mRNA, complete cds"		BG082209	Mm.6958	Signal Transduction
2030 2031	H3073G07		Mus musculus mRNA for granzyme M, complete cds	BG082217	Mm.22302	Mm.22302 Energy/Metabolism
2032 2033	H3074A02 inhibitor	of apoptosis protein 1	Mus musculus inhibitor of apoptosis protein 1 mRNA, complete cds	BG069214	Mm.2026	Apoptosis
2034 2035	NIMA (ne H3074A03 expressed	ever in mitosis gene a)-related kinase 7 (Nek7),	Mus musculus NIMA (never in mitosis gene a)-related expressed kinase 7 (Nek7), mRNA		Mm.143817 Cell Cycle	Cell Cycle
2036 2037	H3074C12 dUTPase		Mus musculus dUTPase mRNA, complete eds	BG082266		Energy/Metabolism

2038	"Mus m type 2A H3074F08 mRNA.	"Mus musculus protein phosphatase type 2A catalytic subunit alpha isoform mRNA. complete cds"		BG082298	Мт.11711	Mm.11711 Signal Transduction
2040	H3074G02	is 40 kDa protein kinase related tK2	H.sapiens 40 kDa protein kinase related to rat BRK2		Mm.1700	
2042	H3074H09	e II protein	Homo sapiens mRNA for NAALADase Il protein	BG082322	Mm.26132	Energy/Metabolism
2044 2045	H3075A04	H3075A04Mm.26128			Mm.26128	Mm.26128 Energy/Metabolism
2046	H3075A09	esterase-22=endoplasmic reticulum- targeting protein of beta-glucuronidase H3075A09[mice,, 2022 nt]	esterase-22=endoplasmic reticulum- targeting protein of beta-glucuronidase [mice, mRNA, 2022 nt]	BG069315	Mm.29110	Mm.29110 Energy/Metabolism
2048 2049	H3075F01	cinyl-CoA synthetase), partial cds	Mus musculus ATP-specific succinyl- CoA synthetase beta subunit (Scs) mRNA, partial cds	BG069442	Mm.19154	Mm.19154 Energy/Metabolism
2050 2051	H3075F08		IA for F1F0-ATP	BG082389	Mm.14663	Mm.14663 Energy/Metabolism
2052 2053	H3075F11	"Mus musculus heat shock protein 20- H3075F11 like protein mRNA, complete cds"		BG069463	Mm.21549	Mm.21549 Heat Shock/Stress
2054 2055	molybde proteins H3075G06 s, partial	num cofactor synthesis-step 1 A and B splice type I (Mocs1) cds, alternatively spliced	Mus musculus molybdenum cofactor synthesis-step 1 proteins A and B splice type I (Mocs1) mRNAs, partial cds, alternatively spliced	BG082399	Mm.36112	Mm.36112 Energy/Metabolism
2056 2057	peptidyl H3075H02 (Pdi2),	arginine deiminase, type Π	Mus musculus peptidyl arginine deiminase, type II (Pdi2), mRNA	BG082407	Мт.2296	Energy/Metabolism
2058 2059	H3076B05	"Mus musculus Rho interacting protein H3076B05[2 (Rhoip2-pending), mRNA"		BG069493	Mm.4620	Signal Transduction

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2060	H3076C12	fatty-acid-Coenzyme A ligase, long-	Homo sapiens fatty-acid-Coenzyme A ligase, long-chain 4 (FACLA), mRNA	BG069344	Мт.12166	Mm.12166 Energy/Metabolism
	H3076F04	"AC005290, complete sequence [Mus	F	BG069372	Mm.1967	Signal Transduction
21	H3076F12	"Homo sapiens regulator of G protein signaling RGS12 (RGS) mRNA, complete cds"		BG082484 Mm.31378	Mm.31378	Signal Transduction
2066	"Mus mus activator	"Mus mi activator		BG069385	Mm.21935	BG069385 Mm.21935 Signal Transduction
2068	similar tr	o argininosuccinate lyase (H. (LOC63402),	Homo sapiens similar to argininosuccinate lyase (H. sapiens) (LOC63402), mRNA	BG082495	Mm.23869	Mm.23869 Energy/Metabolism
2070	"Mus I H3076H06(Ak4),	1 5 7		BG082501	Mm.142051	Mm.142051 Signal Transduction
2072		1 😑		BG069401	Mm.28881	Signal Transduction
2073 2074	H3076H102, polyg	Deptide (CsnK2a2), mKVA- arrier family 12, member 2	Mus musculus solute carrier family 12, member 2 (Sic12a2), mRNA	1	Mm.4168	Energy/Metabolism
2076	H3077C0]	Homo sapiens X28 region near ALD X28 region near ALD locus containing locus containing dual specificity dual specificity phosphatase 9 (DUSP9), phosphatase 9 (DUSP9), ribosomal ribosomal protein L18a (RPL18a), Ca2+/Calmodulin-dependent protein Ca2+/Calmodulin-dependent protein H3077C01 kinase I (CAMKI), creatine transporter (kinase I (CAMKI), creatine transporter (BG069516 Mm.28056 Energy/Metabolism	Homo sapiens X28 region near ALD locus containing dual specificity phosphatase 9 (DUSP9), ribosomal protein L18a (RPL18a), Ca2+/Calmodulin-dependent protein Kinase I (CAMKI), creatine transporter (Kinase I (CAMKI), creatine transporter (BG069516	Mm.28056	Energy/Metabolism

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Mm.26909 Energy/Metabolism	70	Cell Cvole	BG082674 Mm.26207 Energy/Metabolism	Wm.26207 Energy/Merabolism	Energy/Metabolism	Signal Transduction	Heat Shock/Stress	Mm.22675 Enerov/Metaholism
Mm.26909	Mm.30718	Mm.143817 Cell Cycle	Mm.26207	Mm.26207	Mm.4168			Mm.22675
BG082574			BG082674	BG082676	BG069726	BG069727	BG069733	BG069739
Homo sapiens ATPase, H(+)- transporting, lysosomal, noncatalytic accessory protein 1B (ATP6N1B), mRNA	Homo sapiens hypothetical protein FLJ10726 (FLJ10726), mRNA	Mus musculus NIMA (never in mitosis gene a)-related expressed kinase 7 (Nek7), mRNA	Rat L-gulono-gamma-lactone oxidase mRNA, complete cds	Homo sapiens TRPM-2, cytosolic epoxide hydrolase, nicotinic acetylcholine receptor alpha2 subunit, and focal adhesion kinase genes, complete cds, complete sequence	Mus musculus solute carrier family 12, member 2 (Slc12a2), mRNA			Homo sapiens cDNA: FLJ22083 fis, clone HEP14459, highly similar to HUM3H3M Homo sapiens 3-hydroxy-3-methylglutaryl coenzyme A synthase mRNA
ATPase, H(+)-transporting, lysosomal, noncatalytic accessory protein 1B H3077F08 (ATP6N1B),	hypothetical protein FLJ10726 H3077H01 (FLJ10726),	NIMA (never in mitosis gene a)-related H3078G02 expressed kinase 7 (Nek7),	H3078H09 Rat L-gulono-gamma-lactone oxidase	TRPM-2, cytosolic epoxide hydrolase, nicotinic acetylcholine receptor alpha2 subunit, and focal adhesion kinase H3078H11 genes, sequence	solute carrier family 12, member 2 H3079D11 (Slc12a2),	nusculus phospholipase D3 mRNA"	"Mus musculus interferon regulatory factor 6 (Irf6), mRNA"	LJ22083 fis, clone HEP14459, milar to HUM3H3M 3- 3-methylglutaryl coenzyme A
H3077F08	H3077H0	H3078G0	H3078H09	H3078H11	H3079D11	"Mus 1 H3079D12 (Pld3),	"Mus m H3079E06 factor 6	cDNA: F highly si hydroxy- H3079F01 synthase
2078 2079	2080 2081	2082 2083	2084 2085	2086 2087	2088 2089	2090 2091	2092 2093	2094 2095

2096	H3070E05	"Mus musculus calcyclin binding	Щ	BG069742 I	Mm.10702	Signal Transduction
	cystathion major and H3079G04 products			BG082744	Mm.24225	Energy/Metabolism
	"Mus mu	"Mus musculus interferon regulatory factor 3 (frf3), mRNA"		BG069754	Мт.3960	Heat Shock/Stress
2102	H3079G12	KFZp727E011 (from clone 27E011); partial cds	Homo sapiens mRNA; cDNA DKFZp727E011 (from clone DKFZp727E011); partial cds	BG082752	Mm.24225	Energy/Metabolism
2104	H3079H11 M.muscu		M.musculus RPS3a gene	BG069771	Mm.6957	Apoptosis
2106	"Homo ("Homo (H3080A09)	"Homo sapiens similar to cullin 5 (H. 9)saniens) (LOC63330), mRNA"		BG069781	Mm.21710	Mm.21710 Signal Transduction
2108	TF-1 cel	ll apoptosis related protein-15	Mus musculus TF-1 cell apoptosis related protein-15 (Tfar15), mRNA	BG069793	Mm.28441 Apoptosis	Apoptosis
2110	113080CT	sculus ribonuclease P2		BG069849	Mm.12440	Mm.12440 Energy/Metabolism
2112	H3080G1	"Mus musculus adenomatosis polyposis		BG069851	Mm.7883	Signal Transduction
2114	H3080H1	Mus musculus RNA guanylyltransferase H3080H10 and 5-phosphatase (Rngtt), mRNA		BG069862	Mm.26153	Energy/Metabolism
2116 2117	H3081A1	fatty acid-Coenzyme A ligase, long H3081A12 chain 4 (Facl4),	Mus musculus fatty acid-Coenzyme A ligase, long chain 4 (Facl4), mRNA	BG069876	Mm.14368	Mm.143689 Energy/Metabolism

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2118	H3081D02	Mus musculus Bcl-2-related ovarian killer protein-like- killer protein-like-pending (Bokl-H3081D02)pending (Bokl-pending),	Mus musculus Bel-2-related ovarian killer protein-like-pending (Bokl- pending), mRNA	BG082842 Mm.3295		Apoptosis
2120 2121	H3081E08	TNF-alpha converting enzyme H3081E08 precursor, , alternatively spliced	Mus musculus TNF-alpha converting enzyme precursor, mRNA, alternatively spliced, complete cds	BG082857	Mm.27681	Apoptosis
2122 2123	H3081H02		Mus musculus aldo-keto reductase AKR1C13 (Akr1c13), mRNA	BG082884	Mm.27447	Energy/Metabolism
2124 2125	H3082B04	Mus musculus inducible 6- phosphofructo-2-kinase mRNA, H3082B04 complete cds		BG070045	Mm.19669	Mm.19669 Energy/Metabolism
2126 2127	Mus mu H3082C01 1-5 (Spi	Mus musculus serine protease inhibitor 1-5 (Spi1-5), mRNA		BG070053	Mm.89871	BG070053 Mm.89871 Energy/Metabolism
2128 2129	H3082E03	Mus musculus programmed cell death 8 (apoptosis inducing factor) (Pdcd8), H3082E03 mRNA		BG082930	Mm.30050	Apoptosis
2130 2131	H3082E10	H.sapiens mitogen inducible gene mig- H3082B10 2, complete CDS			Mm.29842	Signal Transduction
2132 2133	H3082F03	M.musculus (balb-c) gene for H3082F03 sphingomyelin phosphodiesterase		BG082939		Energy/Metabolism
2134 2135	H3083C01	Homo sapiens kinesin-like 5 (mitotic H3083C01 kinesin-like protein 1) (KNSL5), mRNA			Mm.28386 Cell Cycle	Cell Cycle
2136 2137	Н3083D06	Mus musculus Bcl2-associated H3083D06 athanogene 1 (Bag1), mRNA	·	BG069990	Мт.688	Apoptosis
2138 2139	Н3083D09	Homo sapiens cDNA FLJ14310 fis, H3083D09 clone PLACE3000271		BG069993	Mm.7156	Energy/Metabolism

2140	"Mus mu H3083E02 enzyme ("Mus musculus beta-site APP cleaving enzyme (Bace), mRNA"		BG069998	Mm.24044	Mm.24044 Signal Transduction
	H3083H02			BG070116	Mm.5305	Signal Transduction
	Mus mus necrosis	Mus musculus mRNA for tumor necrosis factor alpha converting enzyme (TACF/ADAM17), complete cds		BG070119	Mm.27681	Energy/Metabolism
2145	H3084E05	ne S-transferase, mu 5 (Gstm5),	one S-transferase,	BG070174 Mm.667	Mm.667	Energy/Metabolism
2147	H3084F02	H3084F02 Mm.24864			Mm.24864	Mm.24864 Energy/Metabolism
2149	H3084G0	Mouse c-fos gene; cellular homolog to H3084G03 viral oncogene		BG070196 Mm.5043	Mm.5043	Signal Transduction
2151	M.mu H3085B01 alpha	M.musculus mRNA for casein kinase I- I alpha		BG083155	Mm.43737	BG083155 Mm.43737 Signal Transduction
2153	Mus mu associat H3085C02 mRNA	Mus musculus Traf and Tnf receptor associated protein (Trrap-pending),		BG070243	Mm.155228 Apoptosis	Apoptosis
2155	H3085C0	Mus musculus solute carrier family 12, H3085C04 member 2 (Slc12a2), mRNA		, BG070245	Mm.4168	Energy/Metabolism
2157	H3085D1	Mus musculus guanine deaminase		BG083182	Mm.25912	Energy/Metabolism
2159	Mov H3085G03 end	Mouse cytochrome beta-558 mRNA, 3		BG083209	Mm.448	Energy/Metabolism
2161	H3086E09	Homo sapiens BCL2-associated H3086E09 athanogene 5 (BAG5), mRNA		BG070363	BG070363 Mm.44239 Apoptosis	Apoptosis

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Mm.28058 Energy/Metabolism	Signal Transduction	Energy/Metabolism	Signal Transduction	Mm.104491 Heat Shock/Stress	Energy/Metabolism	Signal Transduction	Energy/Metabolism	Apoptosis	Signal Transduction	
Mm.28058	Mm.22261	Mm.7612		Mm.104491		Mm.1221	Mm.7446	BG083524 Mm.153450 Apoptosis	Mm.5262	
BG083295	BG083317	BG070413	BG083351	BG070480	BG083405 Mm.1776	BG083409	BG070501	BG083524	BG070547	
		Mouse uterine lactotransferrin mRNA			Human mRNA for acetoacetyl- coenzyme A thiolase (EC 2.3.1.9)		Mus musculus glutathione S-transferase mu 2 (Gstm2), mRNA	Mus musculus CPP32 apoptotic protease mRNA, partial cds		
B.taurus CI-SGDH mRNA for H3086F10 lubiquinone oxidoreductase complex	"Mus musculus mRNA for Rho guanine nucleotide-exchange factor, splice variant NET1A"	uterine lactotransferrin	"Mus musculus rho GTP ase activating protein 6 (Arhgap6), mRNA"	"Mus musculus heat shock protein 030 H3087G09(Hsp030), mRNA"	me A thiolase (EC	"Mus musculus calcium binding protein A6 (calcyclin) (S100a6), mRNA"	Mus musculus glutathione S-transferase, mu 2 (Gstm2), mu 2 (Gstm2), mRNA	H3088C12 CPP32 apoptotic protease, partial cds	"Mus musculus choline kinase (Chk), mRNA"	
H3086F10	"Mus n nucleot H3086H09 variant	H3087A12 uterine	"Mus n H3087C10 protein	H3087G09	acetoac H3087H05[2.3.1.9]	"Mus m H3087H09 A6 (cal	H3088A07	H3088C12	"Mus m H3088E07 mRNA'	
2163 2164	2165 2166	2167 2168	2169 2170	2171 2172	2173 2174	2175 2176	21 <i>77</i> 21 <i>7</i> 8	21 <i>7</i> 9 2180	2181 2182	

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2185	- + 1	Mus musculus ATPase, H+ transporting, lysosomal (vacuolar proton pump), beta 56/58 kDa, isoform	Á	BG083588 N	Mm.10727	Energy/Metabolism	
	114000	Mus musculus ATP-binding cassette		BG083593 N	лт.21629 Л	Mm.21629 Energy/Metabolism	
	3089A11	"Mus musculus protein kinase Chk2	M .	BG083482	Vm.126031	Mm.126031Signal Transduction	
2190 H 2191 H	H3089F09 (Kaccost	Mus musculus zinc transporter like 1	Д	BG083497	Mm.28851	Energy/Metabolism	
		Mouse normal c-myc gene and translocated homologue from J558			74 m	Girnel Transduction	
	13089H11	H3089H11 plasmocytoma cells (cDNA sequence)		BGU&52390	William Street	organical removements	Τ
2196 2196 7197	H3090B01 related-1	mRNA, complete cds" NA glycosylase gene,	Mus musculus uracil-DNA glycosylase	3G083610	Mm.15185	BG083610 Mm.15185 Signal Hansunction	
	H3090B05	те	gene, complete cds, alternatively spinced BCO 70097. Mus musculus omithine decarboxylase BG070710 antizyme inhibitor (Oazi), mRNA		Mm.6775	Energy/Metabolism	T
2201 2202 1	N-termi H3090C10 (Ntan1)	nal asparagine amidohydrolase gene	gine	BG083629	Mm.22635	Mm.22635 Energy/Metabolism	

Signal Transduction	Mm.42205 Energy/Metabolism	7 Mm.133949 Signal Transduction	Apoptosis	Energy/Metabolism	Energy/Metabolism	Energy/Metabolism	Signal Transduction	Mm.154427 Energy/Metabolism	Energy/Metabolism	Apoptosis
Mm.4358	Mm.42205	Mm.133949	BG070891 Mm.22585 Apoptosis	Мт.41722	Mm.22635	Mm.1056	Mm.6793	Mm.154427	Mm.43415	BG070820 Mm.22279 Apoptosis
BG083632	BG070787	BG083726	BG070891	BG070894	BG070912	BG070928	BG070933	BG070934	BG083822	BG070820
	Mus musculus partial gene encoding deoxyribonuclease I and 5 UTR		Mus musculus tumor necrosis factor (ligand) superfamily, member 19 (Tnfsf19-pending), mRNA	Mus musculus 5(3)- deoxyribonucleotidase (Dnt-pending), mRNA	Mus musculus N-terminal Asn amidase (Wtan1), mRNA	Mus musculus solute carrier family 1, member 7 (Slc1a7), mRNA		Rattus norvegicus deiodinase, iodothyronine type III (Dio3), mRNA	Mus musculus Balb/c cytochrome c oxidase subunit VIaL mRNA, complete cds	Mus musculus caspase 8 associated protein 2 (Casp8ap2), mRNA
"Mus musculus mitogen activated protein kinase kinase kinase 12 H3090D01 (Map3k12), mRNA"	partial gene encoding H3091B08 deoxyribonuclease I and 5 UTR	"Mus musculus guanine nucleotide binding protein, alpha 14 (Gna14), mRNA"	tumor necrosis factor (ligand) superfamily, member 19 (Tnfsf19- pending),	5(3)-deoxyribonucleotidase (Dnt-pending),	H3091F08 N-terminal Asn amidase (Ntan1),	solute carrier family 1, member 7 (SIG1a7).	"Mus musculus H-ras gene, partial cds"	deiodinase, iodothyronine type Π	Balb/c cytochrome c oxidase subunit VIaL	caspase 8 associated protein 2 H3092E04 (Casp8ap2),
H3090D01	H3091B08	"Mus m binding H3091D08mRNA"	tumor ne superfam H3091D11 pending)	5(3)-deox H3091E02 pending),	H3091F08	solute c H3091G12(Slc1a7	H3091H05 "Mus m	deiodir H3091H07(Dio3),	Balb/c (H3092E03 VIaL	H3092E04
2203 2204	2205 2206	2207 2208	2209	2211	2213 2214	2215	2217 2218	2219	2221	2223 2224

2225 2226	H3092G01	"Mus musculus p38delta MAP kinase H3092G01 mRNA, complete cds"		BG083840	Mm.27970	BG083840 Mm.27970 Signal Transduction
2227 2228	vacuolar a H3092H01 subunit A	adenosine triphosphatase gene	Mus musculus vacuolar adenosine triphosphatase subunit A gene, complete cds	BG070846 Mm.29771	Mm.29771	Energy/Metabolism
2229 2230	H3092H09	"Mus musculus mitogen-activated protein kinase 8 interacting protein 2 H3092H09(Mapk8ip2), mRNA"		BG083857		Signal Transduction
2231 2232	H3092H10	H3092H10 glycogenin 1 (Gyg1),	Mus musculus glycogenin 1 (Gyg1), mRNA	BG070855	Mm.6375	Energy/Metabolism
2233 2234	H3093B07 chloride	chloride channel 5 (Clon5),	Mus musculus chloride channel 5 (Clcn5), mRNA	BG083875	Mm.104172	Mm.104172Energy/Metabolism
223 5 2236	H3093D09 Tom22		Homo sapiens mRNA for Tom22, complete cds	BG083900 Mm.9326		Energy/Metabolism
_ 2237 2238	H3093F02 uridine m	onophosphate kinase (UMPK),	onophosphate	BG083915		Energy/Metabolism
2239 2240	"Homo se H3093G07 tumor sur	"Homo sapiens Putative prostate cancer tumor suppressor (N33), mRNA"		BG071015	Mm.24156	Signal Transduction
2241 2242	H3093G08	GST-5=glutathione S-transferase-sperm GST-5=glutathione S-transferase-sperm antigen MSAg-5 fusion protein {3 region} [mice, testis, Recombinant region} [mice, testis, Recombinant Recombinant Partial, 860 nt]		BG071016	Mm.11346	Mm.11346 Energy/Metabolism
2243 2244	mannosyl beta-1,2-b acetylgluc H3093H01 (MGAT2)	(alpha-1,6-)-glycoprotein i- iosaminyltransferase	Homo sapiens mannosyl (alpha-1,6-)- glycoprotein beta-1,2-N- acetylglucosaminyltransferase (MGAT2), mRNA	BG084016	Mm.24293	BG084016 Mm.24293 Energy/Metabolism

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Energy/Metabolism	Mm.108678 Energy/Metabolism	Signal Transduction	Mm.29010 Energy/Metabolism	Mm.26092 Energy/Metabolism	Energy/Metabolism		Apoptosis		Energy/Metabolism
	Mm.108678		Мт.29010	Mm.26092	Мт.12983	Mm.29588	BG071119 Mm.17912 Apoptosis		l i
BG084049	BG071067	BG071068 Mm.2344	BG071070		BG083967		BG071119		BG084006 Mm.282
Mus musculus gene for prostacyclin synthase, 5-flanking region and partial cds	Mus musculus cytochrome P450 side chain cleavage enzyme 11a1 H3094D01 (Cyp11a1), mRNA	"Mus musculus GNB1 gene for G protein beta1 subunit, exon 12 and H3094D02 complete cds"	Mus musculus GLUT4 vesicle protein H3094D04 mRNA, partial cds		Homo sapiens cDNA FLJ12225 fis, clone MAMMA1001139, weakly similar to SRE-2 PROTEIN	Homo sapiens mitogen-activated protein kinase kinase kinase kinase 3 H3094G10 (MAP4K3), mRNA	Mus musculus Bcl2-interacting killer- H3094H06 like (Biklk), mRNA	Mus musculus mitogen activated protein kinase kinase kinase 12 (Map3k12), mRNA	Mus musculus glutathione S-transferase H3095E06 like (Gsttl-pending), mRNA
Mus synt H3094C01 cds	H3094D0]	H3094D02	H3094D04	H3094E12	Homo s clone Iv H3094G04 similar	H3094G10	H3094H06	Mus m kinase l H3095D11 mRNA	H3095E06
2245 2246	2247 2248	2249 2250	2251 2252	2253 - 2254	2255 2256	22 <i>57</i> 22 <i>5</i> 8	2259 2260	2261 2262	2263

2264 2265	Homo s H3095F10 protein	Homo sapiens calcineurin binding protein 1 (KIAA0330), mRNA		BG071186	BG071186 Mm.154496	
2266 2267	Rattus no H3095G051, tissue	Rattus norvegicus Fucosidase, alpha-L- 51, tissue (Fuca), mRNA		BG084074	Mm.21580	BG084074 Mm.21580 Energy/Metabolism
2268	Mus m H3095H12 (Car4),	Mus musculus carbonic anhydrase 4 2(Car4), mRNA		BG071207 Mm.1641	Mm.1641	Energy/Metabolism
2269 2270	cytochr H3096A03(Cox4),	ome c oxidase, subunit IV	Mus musculus cytochrome c oxidase, subunit IV (Cox4), mRNA	BG084085	Mm.2136	Energy/Metabolism
2271 2272	H3096D01	"Mus musculus protein phosphatase 4, H3096D01 catalytic subunit (Ppp4c), mRNA"		BG071232	Mm.28082	Signal Transduction
2273 2274	H3096E12	clone TSIP1 p53-induced apoptosis H3096E12 differentially expressed sequence	Mus musculus clone TSIP1 p53- induced apoptosis differentially expressed mRNA sequence	BG084126	BG084126 Mm.142872 Apoptosis	Apoptosis
2275 2276	H3096F11 HNK-1	Ţ),	Homo sapiens HNK-1 sulfotransferase (HNK-1ST), mRNA	BG084134	Мт.26817	BG084134 Mm.26817 Energy/Metabolism
2277 2278	H3097A02 (LOC51	Homo sapiens leucine aminopeptidase (LOC51056), mRNA		BG084150	Mm.34832	BG084150 Mm.34832 Energy/Metabolism
2279 2280	H3097A03	Mus musculus cytochrome c oxidase, H3097A03 subunit IV (Cox4), mRNA		BG084151	Мт.2136	Energy/Metabolism
2281 2282	Homo : H3097A05 mRNA	Homo sapiens sulfite oxidase (SUOX), mRNA		BG084153	Mm.23352	BG084153 Mm.23352 Energy/Metabolism
2283 2284	Mus m aminot H3097B01 mRNA	Mus musculus branched chain aminotransferase 1, cytosolic (Bcat1), mRNA		BG084158 Mm.4606	Mm.4606	Energy/Metabolism

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	Mm.36460 Enerov/Metsholism	Mm.22409 Energy/Metabolism	Hnerm/Metabaliam	Vm 30763 Rusermy Metch of com-	Mm 24916 Simol Targetting	orginal transduction	Mr. 28712 Gignal Itanschiction	Euch By/Metabolism	Signal Transduction Signal Transduction
		_	Mm 4954			Mrs 12145	Mr. 28712	1	
	BG084160		BG084201	BG084205	BG084218	RG084731	B(7)84240		BG084265 Mm.2817 AU041108 Mm.1059
							B.taurus mRNA for PSST subunit of the NADH: ubiquinone oxidoreductase complex		
	Mus musculus solute carrier family 11 (proton-coupled divalent metal ion transporters), member 2 (Slc11a2), mRNA		Mus musculus sialytransferase 8 (alpha-2, 8-sialytransferase) B (Siat8b), mRNA	Homo sapiens FACL5 for fatty acid coenzyme A ligase 5 (LOC51703), mRNA	"Mus musculus retinoblastoma-binding protein 9 (Rbbp9), mRNA."	"Mus musculus retinoblastoma binding protein 4 (Rbbp4), mRNA."	e NADH:	isculus protein tyrosine ase, receptor-type, F interacting binding protein 2 (Ppfibp2),	"Mus musculus mitogen activated protein kinase kinase 1 (Map2k1), mRNA"
	Mus m (protor transpo H3097B03 mRNA	H3097C12	Mus mi (alpha-: H3097F02 mRNA	Homo s coenzyi H3097F06 mRNA	"Mus II H3097G07 protein	"Mus m H3097H08 protein	H3098A081	"Mus m phosphat protein, H3098D02mRNA"	"Mus mu protein 1 H3098D12 mRNA"
	2285 2286	2287 2288	2289 2290	2291 2292	2293 2294	2295 2296	2297 2298	2299 2300	2301 2302

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بلنو	13098E11	"Mus musculus chaperonin subunit 2		BG084277	Ит.40849 Л	BG084277 Mm.40849 Heat Shock/Stress
	H3098H02 carbonyl	C51181),	Homo sapiens carbonyl reductase (LOC51181), mRNA	BG084296		Energy/Metabolism
	5-nucleo H3099A05(NT5B),	tidase (purine), cytosolic type B		BG084308	Мт.29601	Energy/Metabolism
	H3099C01	rrier family 34 (sodium te), member 2 (Sic34a2),	Mus musculus solute carrier family 34 (sodium phosphate), member 2 (Slc34a2), mRNA	BG084323	·	Energy/Metabolism
	И3099D0	t gene	Mouse cytochrome c oxidase Vb subunit gene, complete cds	BG071472	Mm.16769	BG071472 Mm.16769 Energy/Metabolism
	HIZOOORO	cDNA FLJ13488 fis, clone PLACE1003915, weakly similar to PROBABLE ARGINYL-TRNA SYNTHETASE, CYTOPLASMIC (EC.	Homo sapiens cDNA FLJ13488 fis, clone PLACE1003915, weakly similar to PROBABLE ARGINYL-TRNA SYNTHETASE, CYTOPLASMIC (EC 6.1.1.19)	BG084341	Mm.22363	
	H3100C0	P phosphodiesterase delta Pde6d) gene	Mus musculus rod cĠMP phosphodiesterase delta subunit (Pde6d) gene, complete cds	BG084383	Mm.12925	Mm.12925 Energy/Metabolism
	H3100C1	m intermediate/small ance calcium-activated channel, Iy N, member 2 (KCNN2),	Homo sapiens potassium intermediate/small conductance calcium-activated channel, subfamily N, member 2 (KCNN2), mRNA	BG084391	Mm.17734	Energy/Metabolism
	H3100E0	H3100E04 inhibitor of apoptosis protein 1	Mus musculus inhibitor of apoptosis protein 1 mRNA, complete cds	BG084405	Mm.2026	Apoptosis
1	H3101A0		Mus musculus ribosomal protein L23 (Rp[23), mRNA	BG084450	Mm.12144	BG084450 Mm.12144 Energy/Metabolism
2322	H3101A0	8 ribosomai protein L23 (kpl23),	(Kpizz), mikiya	DOLL DODG		_

Energy/Metabolism	Energy/Metabolism	Energy/Metabolism	Mm.29186 Heat Shock/Stress	Mm.38330 Signal Transduction	Energy/Metabolism	Energy/Metabolism	Apoptosis	Signal Transduction	BG071725 Mm.29194 Energy/Metabolism
	Mm.2731	Mm.3786	Mm.29186	Mm.38330		Mm.28056	BG071714 Mm.30236 Apoptosis	Mm.1603	Mm.29194
BG071626 Mm.5289	BG071633	BG071644	AU044106	BG071671	BG071685 Mm.9440	BG071707	 BG071714	BG084566	BG071725
Mus musculus glyceraldehyde-3- phosphate dehydrogenase (Gapd), mRNA	Mus musculus phosphatidylethanolamine N- methyltransferase (Pemt), mRNA	Mus musculus solute carrier family 34 (sodium phosphate), member 2 (Stc34a2), mRNA			Mus musculus phosphatidylserine synthase 1 (Ptdss1), mRNA	Rattus norvegicus choline transporter (CHOT1), mRNA	Mus musculus apoptotic chromatin condensation inducer in the nucleus (Acinus-pending), mRNA		Mus musculus potassium channel modulatory factor DEBT-91 (Debt91), mRNA
glyceraldehyde-3-phosphate	phosphatidylethanolamine N-H3101D05 methyltransferase (Pemt),	solute carrier family 34 (sodium H3101E05 phosphate), member 2 (Slc34a2),	"Mus musculus DNA J protein (Dnaj3),	"Rattus norvegicus mytonic dystrophy kinase-related Cdc42-binding kinase H3101H02 (MRCK) mRNA. complete cds"	rdss1),	1	apoptotic chromatin condensation inducer in the nucleus (Acinus-	"Mus musculus retinoblastoma binding protein 7 (Rbbp7), mRNA"	potassium channel modulatory factor H3102E01 DEBT-91 (Debt91),
H3101C10	H3101D05	H3101E05	"Mus n H3101G05 mRNA	H3101H02	H3102A07	H3102C06 choline	apoptotic inducer ii H3102D02 bending).	"Mus n H3102D09 protein	H3102E01
2323	2325	2327	2329	2330	2332	2334	2336	2338	2340 2341

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Energy/Metabolism	Mm.15622 Energy/Metabolism	Mm.41002 Energy/Metabolism	Energy/Metabolism	Mm.154589 Energy/Metabolism	Heat Shock/Stress	Energy/Metabolism	Apoptosis	Signal Transduction	Apoptosis
Mm.19169 E	Mm.15622 E	Mm.41002 E	ш	Mm.154589 [[]	Mm.29622	Mm.5915	Mm.11526 Apoptosis	Mm.42927	BG071986 Mm.24163 Apoptosis
BG071869	BG071892	BG071908	BG071952	BG084872	BG084881	BG084888	BG084889	BG071976	BG071986
			Pseudomonas aeruginosa PA01, section 341 of 529 of the complete genome	Homo sapiens cDNA FLJ12989 fis, clone NT2RP3000085, moderately similar to ACETYL-/PROPIONYL-COENZYME A CARBOXYLASE ALPHA CHAIN [CONTAINS: BIOTIN CARBOXYLASE (EC 6.3.4.14); BIOTIN CARBOXYL		Homo sapiens CGI-30 protein (LOC51611), mRNA	Mus musculus natural killer cell BY55 precursor (By55-pending), mRNA		Mus musculus masc mRNA for apoptosis-associated speck-like protein containing CARD, complete cds
Mus musculus thioredoxin-like (32kD)	Mouse mRNA for beta-1,4-	Mus musculus ATP-binding cassette protein (Abcc1b) mRNA, partial cds	Pseudomonas aeruginosa PA01, section Pseudomonas aeruginosa PA01, section H3105A09 341 of 529 of the genome 341 of 529 of the complete genome	cDNA FLJ12989 fis, clone NT2RP3000085, moderately similar to ACETYL-/PROPIONYL-COENZYME A CARBOXYLASE ALPHA CHAIN [CONTAINS: BIOTIN CARBOXYLASE (EC 6.3.4.14);	"Mus musculus Cctz-2 gene for chaperonin containing TCP-1 zeta-2	H3105E03 CGL-30 protein (LOCS1611),	natural killer cell BY55 precursor 4/Rv55-nending).	"Mus musculus mRNA for LIMK2b,	masc apoptosis-associated speck-like
11 A POL CTT	H3104D01	Mus mv H3104E06 protein	H3105A09		TOTOLINA TOTOLina TOTOLina TOTOLina TOTOLina TOTOLina TOTOLina TOTolina Tot	H3105B07 subumit,	natural H3105C04/Bv55-	H3105C1(masc at H3105D09protein
2364	2366	2368 2369	2370 2371	2372	2373	2375	2378	2380	2382

					_	
2384 2385	H3105E08	tyrosine 3-monooxygenase/tryptophan r 5-monooxygenase activation protein, r H3105E08 [theta polypeptide (Ywhaq),	Mus musculus tyrosine 3- monooxygenase/tryptophan 5- monooxygenase activation protein, theta polypeptide (Ywhaq), mRNA	3G084914	Vm.14722	BG084914 Mm.14722 Energy/Metabolism
	H2105G04	"Mus musculus mitogen-activated protein kinase kinase kinase kinase kinase 4 https://www.dkd.mRNA."	pad	BG084932	Mm.987	Signal Transduction
	pyrroline H3105G10[P5CR2]	рупоline 5-carboxylate reductase isom	5-carboxylate reductase isom Homo sapiens pyrroline 5-carboxylate reductase isoform (P5CR2), mRNA	BG072021	Mm.29379	Energy/Metabolism
	H3107D11	11				Energy/Metabolism
	H3107E03 H3107E	H3107E03				Energy/Metabolism
	H3107G10	acetylglucosaminidase no disease IIB) (Naglu),	Mus musculus alpha-N- acetylglucosaminidase (Sanfilippo disease IIIB) (Naglu), mRNA	BG072192	Mm.6142	Energy/Metabolism
	010701011	Ę	Homo sapiens carnitine octanovliransferase (COT), mRNA	BG085029	Mm.28197	Mm.28197 Energy/Metabolism
2398	H3108C03 3-5 exon	uclease TREX1	EXI	BG072232	Mm.17632	Mm.17632 Energy/Metabolism
2400	"Homo H3108E02 protein	"Homo sapiens mitogen-activated protein kinase 6 (MAPK6), mRNA"		BG072253	Mm.18856	Mm.18856 Signal Transduction
2402	"Mus m "H3108E05 induced			BG072256	Mm.3903	Signal Transduction
2404 2405	TF-1 ce H3108F08 (Tfar15)	l apoptosis related protein-15	Mus musculus TF-1 cell apoptosis related protein-15 (Tfar15), mRNA	BG085103 Mm.28441 Apoptosis	Mm.28441	Apoptosis

Signal Transduction	Apoptosis	Signal Transduction	BG085195 Mm.42255 Energy/Metabolism	BG085234 Mm.14722 Energy/Metabolism	BG072410 Mm.43737 Signal Transduction	Energy/Metabolism	Signal Transduction	BG085276 Mm.30215 Energy/Metabolism
1	Mm.148395 Apoptosis		Mm.42255	Mm.14722	Mm.43737		Mm.293	Мт.30215
BG072288 Mm.4294	BG085129	BG072299 Mm.1574	BG085195	BG085234	BG072410	BG072453 Mm.9075	BG072475	BG085276
	Mouse laminin B1 mRNA, complete cds	•	Mus musculus ATPase, Ca++ transporting, cardiac muscle, slow twitch 2 (Atp2a2), mRNA	Mus musculus tyrosine 3- monooxygenase/tryptophan 5- monooxygenase activation protein, theta polypeptide (Ywhaq), mRNA		Mus musculus epoxide hydrolase 1, microsomal (Ephx1), mRNA		Mus musculus brain cDNA, clone MNCb-1429, similar to Mus musculus peroxiredoxin V (PrxV) gene
"Mus musculus adenylyl cyclase type 9 H3108H04 mRNA, complete cds"		"Mus musculus receptor (calcitonin) activity modifying protein 2 (Ramp2), mRNA"	Ca++ transporting, cardiac low twitch 2 (Atp2a2),	tophan otein,	"Homo sapiens casein kinase I alpha isoform (CSNK1A1) mRNA, complete cds"	epoxide hydrolase 1, microsomal (Ephx1),	"Mouse brain calmodulin-dependent phosphatase (calcineurin) catalytic subunit mRNA, 3' end"	
H3108H04	H3108H12 laminin	"Mus mu activity n H3109A04mRNA"	H3109F07	H3110B08	"Hoz isofc H3110B12 cds"	H3110F10	H3110H11	H3111A02
2406 2407	2408	2410	2412 2413	2414	2416	2418 2419	2420	2422 2423

2424 2425 H31						
T	11408	"Mus musculus signal transducer and transcription activator 5a (Stat5a) gene, H3111A08 partial cds"	ш	BG085280	, ,	Signal Transduction
	11500	olutathione-S-transferase	Rat Y-b3 glutathione-S-transferase mRNA, complete cds	BG072517	Vim.29640	Mm.29640 Energy/Metabolism
242/ H31 2428 H31	111G06	type II (Arg2),	se type II (Arg2),	BG085317 h	Mm.3506	Energy/Metabolism
	111G12	sporter like 2 (Zatl2)	Mus musculus zinc transporter like 2 (Zntl2) mRNA, complete cds	BG085322 I	Mm.28490	Mm.28490 Energy/Metabolism
	112A04	Mus musculus carnitine H3112A04 palmitoyltransferase 2 (Cpt2), mRNA		BG085333	Mm.29499	Matrix/Structural Proteins
	112805			BG072550	Mm.33981	Heat Shock/Stress
	112B07	H3112B07 Rat mRNA for 3-oxoacyl-CoA thiolase		BG085346	Mm.142498	Mm.142498 Energy/Metabolism
	112F06	"Mus musculus Cetb gene for chaperonin containing TCP-1 beta H3112F06 subunit, complete cds"		BG085384	Mm.3670	Heat Shock/Stress
	1112G09	Mus musculus ATPase, H+ transporting, lysosomal (vacuolar H3112G09[proton pump), 42 kDa (Atp6d), mRNA		BG085399	Mm.1081	Energy/Metabolism
	1113B07	13kDa differentiation-associated protein Homo sapiens 13kDa differentiation- H3113B07(LOC55967),	Homo sapiens 13kDa differentiation-associated protein (LOC55967), mRNA BG085428 Mm.27886 Energy/Metabolism	BG085428	Мт.27886	Energy/Metabolism

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2444 2445	F1F0A7 H3113B08 (Ap5k)	P synthase complex E subunit gene	Mus musculus F1F0ATP synthase complex E subunit (Atp5k) gene, complete cds		Mm.19370	Mm.19370 Energy/Metabolism
2446 2447	"Mus m serine/th H3113E03 mRNA"	"Mus musculus MAP kinase-interacting serine/threonine kinase 2 (Mknk2), mRNA"		BG085457	Mm.6797	Signal Transduction
2448 2449	H3113F04	"Mus musculus kappaB-Ras1 mRNA, H3113F04 complete cds"		BG072675	Mm.25648	Mm.25648 Signal Transduction
2450 2451	ribonuc H3113F06 (RNH),	:lease/angiogenin inhibitor	Homo sapiens ribonuclease/angiogenin inhibitor (RNH), mRNA	BG085471	Mm.6586	Energy/Metabolism
2452 2453	H3113G07	"Homo sapiens phospholipase C, gamma 2 (phosphatidylinositol- H3113G07 specific) (PLCG2), mRNA"		BG085484	Mm.22370	Mm.22370 Signal Transduction
2454 2455	ornithir H3113G08(Oaz3),	te decarboxylase antizyme 3	Mus musculus ornithine decarboxylase antizyme 3 (Oaz3), mRNA	BG085485	Mm.154432	Mm.154432 Energy/Metabolism
2456 2457	H3113H12	"Mus musculus manic fringe homolog H3113H12 (Drosophila) (Mfng), mRNA"	,	BG072704	Mm.517	Signal Transduction
2458 2459	H3114A05	H3114A05 glutathione peroxidase 4 (Gpx4),	Mus musculus glutathione peroxidase 4 (Gpx4), mRNA	BG072706	Mm.2400	Energy/Metabolism
2460 2461	H3114A06	spermidine/spermine N1-acetyl H3114A06(transferase (Sat),	Mus musculus spermidine/spermine N1-acetyl transferase (Sat), mRNA	BG085499	Mm.2734	Energy/Metabolism
2462 2463	H3114C12	cytosolic class 3 aldehyde dehydrogenase (Adh4) gene, 5 flanking H3114C12 region sequence	Mus musculus cytosolic class 3 aldehyde dehydrogenase (Adh4) gene, 5 flanking region sequence	BG085522	Mm.14798	Energy/Metabolism
2464 2465	H3114D10	33a gene				Apoptosis

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Mm.153463 Signal Transduction	Mm.21414 Heat Shock/Stress	Mm.26612 Energy/Metabolism								
Mm.15346	Mm.21414	Mm.26612		Mm.23963						
BG085554	BG072776	BG085572		BG085579	BG085579 BG072801					BG085579 BG072801 BG072817 BG072826
	f mBNA for	nplete cds				Mus musculus clone UWGC:mbac82 from 14D1-D2 (T-Cell Receptor Alpha Locus), complete sequence	VGC:mbac82 Receptor Alpha	VGC:mbac82 Receptor Alpha nce or F1F0-ATP	VGC:mbac82 Receptor Alpha nce or F1F0-ATP	Mus musculus clone UWGC;mbac82 from 14D1-D2 (T-Cell Receptor Alpha Locus), complete sequence Mus musculus mRNA for FIF0-ATP synthase, g subunit Mus musculus mRNA for ATP synthase gamma-subunit precursor, complete cds
	To the state of th	Mannosyltransferase, complete cds				Mus musculus clone UWGC:mbac82 from 14D1-D2 (T-Cell Receptor Alp Locus), complete sequence	culus clone UW D1-D2 (T-Cell]	Mus musculus clone UWGC:mbac82 from 14D1-D2 (T-Cell Receptor Alpi Locus), complete sequence Mus musculus mRNA for F1F0-ATP synthase, g subunit	culus clone UW D1-D2 (T-Cell] complete sequer sculus mRNA f	culus clone UW D1-D2 (T-Cell] complete sequer sculus mRNA fe g subunit sculus mRNA fe
	F	Rattus nor mannosylt				Mus muse from 14I Locus), o				Mus muse from 14I Locus), co Synthase, synthase, Mus muse gamma-s gamma-s gamma-s
Mus musculus interferon gamma	"Mus mus superoxide complete	H3114H06 PIG-M mannosyltransferase	"Mus musculus phospholipase C, della	H3115A05(Plcd), mRNA"	5(Picd), mRNA" Mus musculus S100A9 gene for 7S100A9 protein exons 1-3	S(Picd), mRNA" Mus musculus S100A9 gene for 7S100A9 protein exons 1-3 clone UWGC:mbac82 from 14D1-D2 (T-Cell Receptor Alpha Locus),	S(Plod), mRNA" Mus musculus S100A9 gene for 7S100A9 protein exons 1-3 clone UWGC:mbac32 from 14D1-D2 (T-Cell Receptor Alpha Locus), 2 sequence "Mus musculus protein phosphatase 2A B'alpha3 regulatory subunit mRNA, 1 partial cys"	H3115A05 [Picd), mRNA" Mus musculus S100A9 gene for H3115B07 S100A9 protein exons 1-3 clone UWGC:mbac82 from 14D1-D2 (T-Cell Receptor Alpha Locus), H3115C02 sequence "Mus musculus protein phosphatase 2A B'alpha3 regulatory subunit mRNA, H3115C11 partial cds" H3115D08 F1F0-ATP synthase, g subunit	H3115H05 (Plcd), mRNA" Mus musculus S100A9 gene for Mus musculus S100A9 gene for Clone UWGC:mbac82 from 14D1-D2 (T-Cell Receptor Alpha Locus), (T-Cell Receptor Alpha Locus), H3115C02 sequence "Mus musculus protein phosphatase 2A B'alpha3 regulatory subunit mRNA, H3115D08 F1F0-ATP synthase, g subunit H3115H06 ribosomal protein L29	Harden H
-2		H3114H06]	77011EA 05	MOTICH	H3115B07 S100A9	H3115B07	H3115B07 S100A9 I clone UV (T-Cell R H3115C02 sequence "Mus mus B'alpha3 B'alpha3	H3115B07 H3115C02 H3115C11	H3115B07 H3115C02 H3115C01 H3115D08	H3115B07 H3115C02 H3115C11 H3115H06
2466	2468 2469	2470 2471		2472	2472 2473 2474	2473 2474 2474 2475	2472 2473 2474 2475 2476 2477	2472 2473 2474 2475 2476 2477 2479 2479	2472 2473 2474 2475 2476 2477 2477 2478 2480 2480 2482	2472 2473 2474 2475 2475 2477 2477 2480 2481 2481 2483

Signal Transduction	Energy/Metabolism		Heat Shock/Stress	BG073025 Mm.30072 Energy/Metabolism	Apoptosis	Energy/Metabolism	Energy/Metabolism	Heat Shock/Stress	Energy/Metabolism	BG1085878 Mm 18942 Enerov/Metaholism
		Mm.42203	Mm.6803	Mm.30072	AW548292 Mm.148197Apoptosis	Mm.4084	Mm.258		Mm.424	Mm 18942
BG085699 Mm.44245		.s.	BG072981	BG073025	AW548292	BG085858	BG085863		BG073136	BG085878
		Mus musculus mRNA for kinesin- related mitotic motor protein		Rat glycogen phosphorylase brain isozyme mRNA, 5 end of cds	Mus musculus apoptosis inhibitor bcl-x (bcl-x) gene, exon 3 and complete cds	Mus musculus mastocytoma N- deacetylase/N-sulfotransferase (Mndns), mRNA	Mus musculus H+ ATP synthase subunit c mRNA, complete cds		Mus musculus Na,K-ATPase beta-3 subunit (Atp1b3) gene, exon 7 and complete cds	Mus musculus cytochrome b-561
"Mus musculus adenylate cyclase activating polypeptide 1 receptor 1 H3116E02 (Adcyap1r1), mRNA"	2H3116G12	H3117A12 kinesin-related mitotic motor protein	"Homo sapiens MEBP-1 mRNA for MAPK-ERK binding protein-1, H3117C04 complete cds"	Rat glycogen phosphorylase brain H3117G06isozyme. 5 end of cds	apoptosis inhibitor bcl-x (bcl-x) gene, H3118E12 exon 3 and cds	mastocytoma N-deacetylase/N- H3118H09 sulfotransferase (Mndns),	4H+ ATP synthase subunit c		Na,K-ATPase beta-3 subunit (Atp1b3) 5 gene, exon 7 and cds	1 (Cutsell
H3116E02	H3116G12 H31160	H3117A12	H3117C04	H3117G06	H3118E12	H3118H09	H3119A04H+AT	H3119B04	Na,K-A H3119B06 gene, ex	0000011011
2487 2488	2489	2491	2493	2495	2497	2498	2500	2502 2503	2504 2505	2506

2508	H3119G03	"Mus musculus protein tyrosine H3119G03 phosphatase 4a1 (Ptp4a1), mRNA"	pee	BG073186	Mm.28909	BG073186 Mm.28909 Signal Transduction
2510	glutathio H3119G08 (Gstad).		Mus musculus glutathione S-transferase, alpha 4 (Gsta4), mRNA	BG073190	Мт.2662	Energy/Metabolism
2512	H3119G12	OKFZp564A126 (from clone 564A126); partial cds	SDNA clone al cds	BG085933	Mm.22710	Mm.22710 Energy/Metabolism
2514 2515	Mus alph H3119H03 cds	Mus musculus FVB/N collagen pro- alpha-1 type I chain mRNA, complete cds		BG073196	Мт.22621	Matrix/Structural Proteins
2516 2517	fatty-acid- H3119H04 chain 3 (F	Coenzyme A ligase, long- ACL3),	Homo sapiens fatty-acid-Coenzyme A ligase, long-chain 3 (FACL3), mRNA	BG073197	Mm.27944	Energy/Metabolism
2518 2519	H3119H06	"Homo sapiens casein kinase 1, gamma H3119H062 (CSNK1G2), mRNA"		BG085938 Mm.29873		Signal Transduction
2520	TT3120A16		Mus musculus isocitrate dehydrogenase mRNA, complete cds	BG085953	Mm.2966	Energy/Metabolism
2522 2523	H3120D0	ase 3, mRNA,		BG085978	Mm.27041	Signal Transduction
2524	geranylg H3120F08 (Ggps1),	nyl diphosphate synthase l	Mus musculus geranylgeranyl diphosphate synthase I (Ggps1), mRNA	BG086008 Mm.4076	Mm.4076	Energy/Metabolism
2526 2527	H3120H0	Homo sapiens full length insert cDNA clone ZC24D06 clone ZC24D06	Homo sapiens full length insert cDNA clone ZC24D06	BG086026		Energy/Metabolism
2528 2529	H3121A0	protein phosphatase type-1 catalytic H3121A04[subunit delta isom (PPCS1D)	Homo sapiens protein phosphatase type-1 catalytic subunit delta isoform (PPCS1D) mRNA, complete eds	BG086037 Mm.4572	Mm.4572	Energy/Metabolism

2530 2531	"Mus mu H3121C02 mRNA"	"Mus musculus follistatin-like (Fstl), mRNA"		BG086054	Mm.22763	BG086054 Mm.22763 Signal Transduction
2532 2533	retinal sh dehydrog H3121E04 pending)	retinal short-chain dehydrogenase/reductase 1 (Rsdr1- pending),	Mus musculus retinal short-chain dehydrogenase/reductase 1 (Rsdr1- pending), mRNA	BG086076	Mm.14063	Energy/Metabolism
2534 2535	H3122B07	"Mus musculus heat-responsive protein H3122B0712. (Hrsp12), mRNA"		BG073398	Mm.2607	Heat Shock/Stress
2536 2537	"Mus H3122D07 kDa,	"Mus musculus heat shock protein, 74 kDa, A (Hspa9a), mRNA"		BG073420	Mm.2849	Heat Shock/Stress
2538 2539	ATP H3122E12 gene	syn	Human ATP synthase beta subunit (ATPSB) gene, complete eds	BG073436	Mm.17869	BG073436 Mm.17869 Energy/Metabolism
2540 2541	H3122F01	synthase beta subunit (ATPSB)	bunit	BG073437	Мт.17869	Mm.17869 Energy/Metabolism
2542 2543	ATP-binc H3122F04 (ALD), n	ling cassette, sub-family D tember 4 (Abcd4),	Mus musculus ATP-binding cassette, sub-family D (ALD), member 4 (Abcd4), mRNA	BG086172	Mm.87806	Mm.87806 Energy/Metabolism
2544 2545	H3122F12	H3122F12 sterol-carrier protein X	Mus musculus sterol-carrier protein X mRNA, complete cds	BG086180	Mm.1779	Matrix/Structural Proteins
2546 2547	H3122G01 sterol-car	rier protein X	Mus musculus sterol-carrier protein X mRNA, complete cds	BG086181	Mm.1779	Matrix/Structural Proteins
2548 2549	H3122H03	Mus musculus brain cDNA, clone MNCb-0663, similar to Mus musculus brain cDNA, clone MNCb-0663, similar augmenter of liver regeneration (Alr) mRNA	usculus brain cDNA, clone -0663, similar to Mus musculus nter of liver regeneration (Alr)	BG086194	Mm.28124	
2550 2551	"Mu bind H3123C04 cds"	"Mus musculus GNB3 gene for GTP-binding protein beta3 subunit, complete cds"		BG073494	Mm.35708	Mm.35708 Signal Transduction

2552	BCL2/ad	enovirus B1B 19kD-interacting (BNIP1).		BG073508 Mm.21795 Apoptosis	Mm.21795	Apoptosis
	mannosi acetylglu H3123G10(Mgat1),	le cosaminyltransferase 1	Mus musculus mannoside acetylglucosaminyltransferase 1 (Mgat1), mRNA	BG073537	Mm.2672	Energy/Metabolism
	"Homo	"Homo sapiens tumor protein p53-		BG086271	Mm.23595	Signal Transduction
2558	novel cell	ill death-regulatory protein 9 (Grim19)	Mus musculus novel cell death- regulatory protein GRIM19 (Grim19) mRNA, complete cds	BG086273	Mm.21162 Apoptosis	Apoptosis
2560	"Mus m" H3124B02 hinding	"Mus musculus cold inducible RNA-		BG086286	Mm.17898	BG086286 Mm.17898 Heat Shock/Stress
2562	H3124D02	latase (Fech),	Mus musculus ferrochelatase (Fech), mRNA	BG086309	Мт.1070	Energy/Metabolism
2564	H3124D04 caspase 9	(caspase 9 (Casp9),	Mus musculus caspase 9 (Casp9), mRNA	BG086310 Mm.102926 Apoptosis	Mm.102926	Apoptosis
2566 2567	DNA fra H3124D06 subunit	agmentation factor, 40 kD, beta (Dfb),	Mus musculus DNA fragmentation factor, 40 kD, beta subunit (Dffb), mRNA	BG086312	Mm.38853	Apoptosis
2568	mitotic a H3124D11 (Mad111	mitotic arrest deficient 1-like 1 [(Mad111),	Mus musculus mitotic arrest deficient 1-like 1 (Mad111), mRNA		Mm.27250 Cell Cycle	Cell Cycle
2570 2571	hepa: H3124E02 gene	tocyte growth factor-like protein	Mouse hepatocyte growth factor-like protein gene, complete cds	BG073592	Mm.21989	Mm.21989 Energy/Metabolism
2572 2573	micro H3124F04 (Gst),	microsomal glutathione S-transferase (Gst),	Mus musculus microsomal glutathione S-transferase (Gst), mRNA	BG086330	Mm.14796	BG086330 Mm.14796 Energy/Metabolism

Energy/Metabolism	Signal Transduction	Energy/Metabolism	Mm.75856 Heat Shock/Stress	Energy/Metabolism	Energy/Metabolism	Heat Shock/Stress	Mm.142740 Energy/Metabolism	Energy/Metabolism
Mm.8688		Mm.7156	Mm.75856		Mm.1457	Mm.4908	Mm.142740	Mm.1275
BG086348	BG086363	BG073718	BG086370	BG086378	BG086380	BG086389	BG086391	BG086400 Mm.1275
Homo sapiens NADH dehydrogenase (ubiquinone) 1 beta subcomplex, 10 (22kD, PDSW) (NDUFB10), mRNA		Mus musculus glutathione peroxidase 3 (Gpx3), mRNA		Homo sapiens ribonuclease III (RN3) mRNA, complete cds	Mus musculus major histocompatibility locus class III regions Hsc70t gene, partial cds; smRNP, G7A, NG23, MutS homolog, CLCP, NG24, NG25, and NG26 genes, complete cds; and unknown genes		Homo sapiens cytosolic acyl coenzyme A thioester hydrolase (HBACH), mRNA	Mus musculus thioredoxin (Txn), mRNA
NADH dehydrogenase (ubiquinone) 1 beta subcomplex, 10 (22kD, PDSW) H3124H01 (NDUFB10),	"Mus musculus glycogen synthase kinase 3 beta (Gsk3b), mRNA"	H3125A09 glutathione peroxidase 3 (Gpx3),	"Homo sapiens heat shock transcription factor 2 binding protein (HSF2BP), mRNA"	ribonuclease III (RN3)	major histocompatibility locus class III regions Hsc70t gene, partial cds; smRNP, G7A, NG23, MutS homolog, CLCP, NG24, NG25, and NG26 genes; H3125C10 and unknown genes	"Homo sapiens suppression of tumorigenicity 13 (colon carcinoma) (Hsp70-interacting protein) (ST13), mRNA"	cytosolic acyl coenzyme A thioester H3125E01 hydrolase (HBACH),	H3125F03 thioredoxin (Txn),
H3124H01	"Mus m H3125A06 kinase 3	H3125A09	"Homo factor 2 H3125B05 mRNA"	H3125C08 ribonucl	H3125C10	"Homo : tumorigi (Hsp70- H3125D09 mRNA"	H3125E01	H3125F03
2574 2575	2576 2577	2578 2579	2580 2581	2582 2583	2584 2585	2586 2587	2588 2589	2590 2591

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Matrix/Structural Proteins	Apoptosis	Mm.10091 Signal Transduction	Energy/Metabolism	Energy/Metabolism	Mm.43660 Energy/Metabolism	Mm.41565 Energy/Metabolism	Mm.14064 Energy/Metabolism	Energy/Metabolism	Energy/Metabolism
Mm.27393	Mm.16228 Apoptosis	Mm.10091	Mm.830	Mm.3553	Mm.43660	Mm.41565	Mm.14064	Mm.77432	Mm.7248
		BG086437	BG073636 Mm.830	BG086479	BG086559	BG086562	BG086604	BG086605	BG086676 Mm.7248
Homo sapiens protein translocase, JM26 protein, UDP-galactose translocator, pim-2 protooncogene homolog pim-2h, and shal-type potassium channel genes, complete eds; JM12 protein and transcription	M.musculus RPS3a gene		Mus musculus protease (prosome, macropain) 28 subunit, alpha (Psme1), mRNA	Mus musculus spermine synthase (Sms), mRNA	Mus musculus palmitoyl-protein thioesterase (Ppt), mRNA	Homo sapiens cDNA FLJ13974 fis, clone Y79AA1001581, weakly similar to ACETYL-COENZYME A SYNTHETASE (EC 6.2.1.1)	Mouse s-adenosylmethionine synthetase mRNA	Mus musculus thioredoxin interacting factor (Vdup1) gene, complete cds	Mus musculus solute carrier family 4 (anion exchanger), member 1 (Slo4a1), mRNA
Protein translocase, JM26 protein, UDP- JM26 protein, UDP-galactose galactose translocator, pim-2 protooncogene homolog pim-2h, and shal-type potassium channel genes; potassium channel genes; IM12 protein and transcription		"Mus musculus protein kinase inhibitor, H3126B12 eamma (Pkig), mRNA"	ropain) 28	(6	palmitoyl-protein thioesterase (Ppt),	cDNA FLJ13974 fis, clone Y79AA1001581, weakly similar to ACETYL-COENZYME A	H3128D01 s-adenosylmethionine synthetase	thioredoxin interacting factor (Vdup1)	solute carrier family 4 (anion H3129C03 exchanger), member 1 (Slc4a1),
H3195B07	H3125H05	H3126B12	C173106713	H3176F10	H3127G05 nalmitov		H3128D01	thior H3128D02 gene	H3129C03
2592	2594 2595	2596	2598	2600	2602	2604	2606	2608	2610

2612	H3129C06	protein-L-isoaspartate (D-aspartate) O-H3129C06 methyltransferase 1 (Pcmt1),	Mus musculus protein-L-isoaspartate (D-aspartate) O-methyltransferase 1 (Pcmt1), mRNA	BG086679	Mm.25169	Mm.25169 Energy/Metabolism
2614	"Homo suppres	"Homo sapiens candidate tumor suppressor p33 ING1 homolog (LOC51147), mRNA"		BG086692	Mm.43482	Signal Transduction
2616 2617	H3129E06 OXA1I	OXA1L gene sequence	Homo sapiens OXA1L gene complete sequence	BG086699	Mm.142455 Apoptosis	Apoptosis
2618 2619	H3129E10	"Mus musculus Pro-rich, PH, SH2 domain-containing signaling mediator H3129E10 (PSM) mRNA, complete cds"		BG086702	Mm.8538	Signal Transduction
2620	H3129E12	Mus musculus IFN alpha-treated H3129E12 embryonic fibroblast mRNA		BG086704	Mm.23836	BG086704 Mm.23836 Heat Shock/Stress
2622	H3129F04	clone TA-9 ATP synthase b chain H3129F04 homolog, partial cds	Mus musculus clone TA-9 ATP synthase b chain homolog mRNA, partial cds		Mm.30112	Mm.30112 Energy/Metabolism
2624 2625	H3129G02	serine palmitoyltransferase, long chain H3129G02base subunit 1 (Sptlc1),	Mus musculus serine palmitoyltransferase, long chain base subunit 1 (Sptlc1), mRNA	BG074002	Mm.6505	Energy/Metabolism
2626 2627	peroxi H3129G06(Pex3)	peroxisomal assembly protein PEX3P (Pex3) gene, promoter and exon 1	Mus musculus peroxisomal assembly protein PEX3P (Pex3) gene, promoter and exon 1	BG086720		Energy/Metabolism
2628 2629	H3129H03	hydroxylacyl-Coenzyme A H3129H03 dehydrogenase (Hadh),	Mus musculus hydroxylacyl-Coenzyme A dehydrogenase (Hadh), mRNA	BG086728	Mm.2491	Energy/Metabolism
2630 2631	H3129H05K-CI co	K-Cl cotransporter KCC3	Homo sapiens K-Cl cotransporter KCC3 mRNA, complete cds	BG086729	Mm.21915	Mm.21915 Energy/Metabolism
2632 2633	H3130B10	H3130B10 SERCA2 gene, alternative transcripts	Mus musculus SERCA2 gene, alternative transcripts	BG074044	Mm.42255	Mm.42255 Energy/Metabolism

2634	H3130B11	"Mouse Cu-Zn superoxide dismutase H3130B11 mRNA, complete cds"		BG074045 Mm.5274	,	Heat Shock/Stress
2636 2637	"Mus m mRNA, H3130C02 spliced"	"Mus musculus calcineurin inhibitor mRNA, complete cds, alternatively spliced"		BG086761	Mm.56	Signal Transduction
2638 2639	H3130D06	TPase beta subunit (ATP1B) ons 1 and 2	Human Na,K-ATPase beta subunit (ATP1B) gene, exons 1 and 2	BG086774		Energy/Metabolism
2640 2641	H3130D08	togenic oxidase	Rattus norvegicus NADH/NADPH mitogenic oxidase subunit p65-mox mRNA, complete cds		Мт.2866	
2642 2643	H3130H07	hyiglutaryi-Coenzyme iHS			Mm.10633	
2644 2645	"Mus H3131A03 kDa 1	"Mus musculus heat shock protein, 86 kDa 1 (Hsp86-1), mRNA"		BG074109	Mm.1843	Heat Shock/Stress
2646 2647	H3131A05	nit precursor	Mus musculus mRNA for ATP synthase gamma-subunit precursor, complete cds	BG074111	Mm.12677	Mm.12677 Energy/Metabolism
2648 2649	H3131A07	enoyl coenzyme A hydratase 1, H3131A07[peroxisomal (Ech1),	Mus musculus enoyl coenzyme A hydratase 1, peroxisomal (Ech1), mRNA	BG074113 Mm.2112	Mm.2112	Energy/Metabolism
2650 2651	H3131A12	ated gene)	Mouse MA-3 (apoptosis-related gene) mRNA, complete cds	BG086831	Mm.26035	Apoptosis
2652 2653	H3131B04		arrier protein X	BG086835	Mm.1779	Matrix/Structural Proteins
2654 2655	H3131C10	gosaccharide- e (Ddost),	yl-di- de-protein st), mRNA	BG074138 Mm.7236		Energy/Metabolism

Signal Transduction	Mm.734 Signal Transduction Mm.23923 Signal Transduction
	BG074245
	"Homo sapiens pleiomorphic adenoma "H3132F06 gene-like 2 (PLAGL2), mRNA" "Homo sapiens ER-associated DNAJ; ER-associated Hsp40 co-chaperone;
-	"Homo sapi 06 gene-like 2 "Homo sapi ER-associat
_	H3132F0
	2664 2665 2666

2678 2679	H3133D04	NADH dehydrogenase (ubiquinone) (H3133D04 flavoprotein 2 (24kD) (NDUFV2),	Homo sapiens NADH dehydrogenase (ubiquinone) flavoprotein 2 (24kD) (NDUFV2), mRNA	BG087002	Mm.2206	Energy/Metabolism
2680 2681	H3133D06				Mm.21535	Mm.21535 Heat Shock/Stress
2682		glycine amidinotransferase (L- arginine:glycine amidinotransferase) (GATM),	Homo sapiens glycine amidinotransferase (L-arginine:glycine amidinotransferase) (GATM), mRNA	BG087011	Mm.29975	Energy/Metabolism
2684	"Mus m H3133H01 (Hsc70)	"Mus musculus heat shock 70 protein (Hsc70) gene, complete cds"		BG087043	Mm.2144	Heat Shock/Stress
2686 2687	H3133H05	ulus arylsulfatase A	M.musculus mRNA for aryisulfatase A BG087046		Mm.620	Energy/Metabolism
2688 2689	voltage- H3133H07 (Vdac3)	dependent anion channel 3	Mus musculus voltage-dependent anion channel 3 (Vdac3), mRNA	BG087048	Mm.133962	BG087048 Mm.133962 Energy/Metabolism
2690 2691	H3134C07	DNA, DLEC1 to ORCTIA gene region, section 1/2 (DLEC1, ORCTL3, H3134C07)ORCTIA genes)	Homo sapiens DNA, DLEC1 to ORCTLA gene region, section 1/2 (DLEC1, ORCTL3, ORCTLA genes, complete cds)	BG087037	Mm.42275	Mm.42275 Energy/Metabolism
2692 2693	GrpE-iil H3134D06 GrpE#2	GrpE-like 2, mitochondrial mt- GrpE#2			Mm.12959	Mm.12959 Heat Shock/Stress
2694 2695	H3135A03	dehydrogenase (ubiquinone) 1 bcomplex, 8 (19kD, PGIV) 48),	Homo sapiens NADH dehydrogenase (ubiquinone) 1 alpha subcomplex, 8 (19kD, PGIV) (NDUFA8), mRNA	BG087056	Mm.19834	BG087056 Mm.19834 Energy/Metabolism
2696 2697	H3135B06	Zp434D175 (from clone 0175); partial cds	Homo sapiens mRNA; cDNA DKFZp434D175 (from clone DKFZp434D175); partial cds	BG087066	Mm.30110	BG087066 Mm.30110 Energy/Metabolism
2698 2699	H3135B10	H3135B10"'Mus musculus PL-6 (Pl6), mRNA"		BG087070 Mm.27312	Mm.27312	Signal Transduction

Energy/Metabolism	Mm.87124 Energy/Metabolism	Heat Shock/Stress	Energy/Metabolism	Energy/Metabolism		Energy/Metabolism	Mm.20897 Energy/Metabolism	Mm.25848 Energy/Metabolism	Apoptosis	BG087171 Mm.24276 Energy/Metabolism		BG087194 Mm.28299 Energy/Metabolism
Mm.2420	Mm.87124		Mm.2623	Mm.4730	Mm.15312	Mm.5731	Мт.20897	Mm.25848	Mm.2159	Mm.24276	Mm.144413	Mm.28299
BG087088			BG087117	BG087120		BG087134	BG087141	BG087153	BG087162	BG087171		BG087194
Mus musculus galactose-1-phosphate uridyltransferase (GALT) gene, complete cds			Mus musculus serine protease inhibitor 3 (Spi3), mRNA	Mus musculus protoporphyrinogen oxidase (Ppox), mRNA	Mus musculus rhodanese (Tst) mRNA, complete cds	Mus musculus D-dopachrome tautomerase gene, complete cds	Mus musculus calcium-sensitive chloride conductance protein-1 (mCLCA1) mRNA, complete cds	Mus musculus branched chain ketoacid dehydrogenase E1, alpha polypeptide (Bckdha), mRNA	Mus musculus BCL2/adenovirus E1B 19 kDa-interacting protein 1, NIP3 (Bnip3), mRNA	Mus musculus serine protease mRNA, complete eds	Mus musculus gene for uncoupling protein-2, complete eds	Homo sapiens guanine monphosphate synthetase (GMPS), mRNA
Mus musculus galactose-1-phos galactose-1-phos galactose-1-phos galactose-1-phosphate uridyltransferase (GALT) gene complete cds			H3135G03 serine protease inhibitor 3 (Spi3),	H3135G07 protoporphyrinogen oxidase (Ppox),	H3135H04rhodanese (Tst)	H3135H10D-dopachrome tautomerase gene	sensitive chloride conductance 1 (mCLCA1)	Mus musculus branched chain ketoaci branched chain ketoaci branched chain ketoaci dehydrogenase E1, alpha polypeptide (Bekdha), (Bekdha), mRNA	BCL2/adenovirus E1B 19 kDa- H3136C07 interacting protein 1, NFP3 (Bnip3),	otcase	H3136E12 gene uncoupling protein-2	monphosphate synthetase
galactos H3135D06(GALT)	H3135F04 Mm.87	H3135F06	H3135G03	H3135G07	H3135H04	H3135H10	calcium H3136A06 protein-	branche H3136B09 E1, alph	H3136C07	H3136D05 serine p	H3136E12	guanine H3136F07 (GMPS)
2700 2701	2702 2703	2704	2705 2706	2707 2708	2709 2710	2711 2712	2713 2714	2715 2716	2717 2718	2719 2720	2721 2722	2723 2724

2725	0130210	sece inhihitor 3 (Sni3)	Mus musculus serine protease inhibitor (Soi3), mRNA	BG087196		Energy/Metabolism
2720 2727 2728	H3136G10(Scine pro solute carr H3136G10(Slc12a2),	ier family 12, member 2	olute carrier family 12, 2a2), mRNA	BG074576 b	Mm.4168	Energy/Metabolism
2729	01H3£1FH	"Mus musculus IkB kinase-beta (Ikkb)		BG087220	Mm.28269	BG087220 Mm.28269 Signal Transduction
2731 2732	H3137A09	e (Pcx),	Mus musculus pyruvate decarboxylase (Pcx), mRNA	BG087228	Mm.1845	Energy/Metabolism
2733	H3137B06 ornithine	at),	Mus musculus ornithine aminotransferase (Oat), mRNA	BG087236	Mm.13694	Mm.13694 Energy/Metabolism
2735	"Homo sa exchange H3137B09 mRNA."	"Homo sapiens Rho guanine nucleotide exchange factor (GEF) 4 (ARHGEF4),		BG087239	Mm.41604	Mm.41604 Signal Transduction
2737	ATP-bin H3137C10(ABC1),	ATP-binding cassette, sub-family A (ABC1), member 3 (ABCA3),	Homo sapiens ATP-binding cassette, sub-family A (ABCI), member 3 (ABCA3), mRNA	BG087249	Mm.38313	Mm.38313 Energy/Metabolism
2739	ATP-bin H3137D07(OABP),		Mus musculus ATP-binding cassette, sub-family B (OABP), member 1 (Abce1), mRNA	BG087256	Mm.5831	Energy/Metabolism
2741	H3137D08 VLCAD	VICAD gene	Mus musculus VLCAD gene	BG074630	Мт.18630	Mm.18630 Energy/Metabolism
2743	solute car H3137D09 exchange		Mus musculus solute carrier family 4 (anion exchanger), member 2 (Slc4a2), mRNA	BG074631	Mm.4580	Energy/Metabolism
2745 2746	H3137E02	NADH dehydrogenase (ubiquinone) Fe- (ubiquinone) Fe-S protein 2 (49kD) (NADH-coenzyme (NADH-coenzyme Q reductase) (NDUFS2), (NDUFS2), mRNA	Homo sapiens NADH dehydrogenase (ubiquinone) Fe-S protein 2 (49kD) (NADH-coenzyme Q reductase) (NDUFS2), mRNA	BG087262	Mm.21669	BG087262 Mm.21669 Energy/Metabolism

Energy/Metabolism	Energy/Metabolism	Signal Transduction	Apoptosis	Energy/Metabolism	Energy/Metabolism	Energy/Metabolism	Energy/Metabolism	Energy/Metabolism	Energy/Metabolism
	Mm.44225		Mm.13437 Apoptosis	Mm.22403			Mm.24457	1	Mm.42253
BG074636 Mm.5053	BG087283	BG087286 Mm.4220	BG087312	BG087325	BG074710 Mm.9870	BG087336 Mm.569	BG087349	BG087356 Mm.1486	BG087364
Mus musculus anion exchanger 3 brain and anion exchanger 3 cardiac isoforms (Slc4a3) gene, complete cds, alternatively spliced	Homo sapiens ribosomal protein L17 isolog (LOC63875), mRNA	-	Mus musculus neuronal apoptosis inhibitory protein 6 (Naip6) gene, complete cds; and Naip3 gene, exons 2- 9 and 11-16	Homo sapiens phosphoglucomutase 1 (PGM1), mRNA	Homo sapiens hypothetical protein FLJ10545 (FLJ10545), mRNA	Mus musculus voltage-dependent anion channel 2 (Vdac2), mRNA	Homo sapiens aldehyde dehydrogenase 5 (ALDH5), mRNA	Mus musculus ADP-ribosylation factor 4 (Arf4), mRNA	Mus musculus solute carrier family 22 (organic cation transporter), member 5 (Slc22a5), mRNA
anion exchanger 3 brain and anion exchanger 3 cardiac isoms (Slc4a3) gene, alternatively spliced	ribosomal protein L17 isolog H3137G05(LOC63875),	Mouse mRNA for protein tyrosine H3137G09phosphatase PTPT9	neuronal apoptosis inhibitory protein 6 (Naip6) gene; and Naip3 gene, exons 2- 9 and 11-16	H3138C04 phosphoglucomutase 1 (PGM1),	hypothetical protein FLJ10545 H3138D04(FLJ10545),	voltage-dependent anion channel 2 (Vdac2),	H3138E10 aldehyde dehydrogenase 5 (ALDH5),	H3138F05 ADP-ribosylation factor 4 (Arf4),	solute carrier family 22 (organic cation H3138G02(transporter), member 5 (Slc22a5),
H3137E03	H3137G05	H3137G09	neurona (Naip6) H3138B029 and 1	H3138C04	H3138D04	voltage- H3138D07 (Vdac2)	H3138E10	H3138F05	H3138G02
2747 2748	2749 2750	2751 2752	2753 2754	2755 2756	2757 2758	2759 2760	2761 2762	2763 2764	2765 2766

2767 2768	alpha gluc H3138G09\subunit ((osidase 2, alpha neutral 32an),	Mus musculus alpha glucosidase 2, alpha neutral subunit (G2an), mRNA	BG087370 Mm.3196	1	Energy/Metabolism
2769	AOLIOGICE	n and Cak cumtherase	Homo sapiens mRNA for long-chain acvl-CoA synthetase, complete cds	BG074754	Mm.28962	BG074754 Mm.28962 Energy/Metabolism
2771	H3138H07		ein	BG087380	Mm.16773	Energy/Metabolism
2773	H3138H10	sin D		BG087383 Mm.2147		Signal Transduction
2775	H3139B06	"Mus musculus protein tyrosine phosphatase, non-receptor type 13 H3139B06[interacting protein (Ptpn13ip), mRNA"		BG087401	Mm.5400	Signal Transduction
2777	H3139C01		Mus musculus ubi-d4/requiem gene, exons 2 through 11 and partial cds	BG087408	Mm.2651	Apoptosis
2779	"Mus m H3139E01 (Hsc70)	at shock 70 protein uplete cds"		BG087426	·	Heat Shock/Stress
2781	H3139F01 sterol O-	-acyltransferase 1 (Soat1),	Mus musculus sterol O-acyltransferase 1 (Soat1), mRNA	BG087432	Mm.28099	Mm.28099 Energy/Metabolism
2783	solute car (sodium/ H3140A021 (SIC9a1	rier family 9 hydrogen exchanger), member),	Mus musculus solute carrier family 9 (sodium/hydrogen exchanger), member 1 (Sic9a1), mRNA	BG087460	Mm.4312	Energy/Metabolism
2785 2786	"Mus m H3140A08 protein	"Mus musculus mitogen activated sprotein kinase 14 (Mapk14), mRNA"		BG087465 Mm.4437	Mm.4437	Signal Transduction

Mm.18522 Energy/Metabolism		Energy/Metabolism	Mm.30145 Energy/Metabolism	Mm.29821 Energy/Metabolism	Mm.26949 Energy/Metabolism	Energy/Metabolism	BG087506 Mm.10600 Energy/Metabolism	Mm.24510 Energy/Metabolism
Mm.18522	Mm.1511		Mm.30145	Mm.29821	Mm.26949	Mm.1484	Mm.10600	Mm.24510
		BG074864 Mm.3479	BG087491	BG087493	BG087498	BG074880	BG087506	BG087669
CPT I=carnitine palmitoyltransferase I {3 region} [mice, IVS, hearts, mRNA Partial, 678 nt, segment 1 of 2]		Homo sapiens clone RP11-486122, complete sequence	Homo sapiens ectonucleotide pyrophosphatase/phosphodiesterase 5 (putative function) (ENPP5), mRNA	Mus musculus peroxiredoxin 3 (Prdx3), mRNA	Homo sapiens electron-transfer- flavoprotein, alpha polypeptide (glutaric aciduria II) (ETFA), mRNA	Mouse protein tyrosine phosphatase (70zshp) mRNA, complete cds	Mus musculus glutamate dehydrogenase (Glud), mRNA	Mus musculus solute carrier family 12, member 7 (Slc12a7), mRNA
CPT I=carnitine palmitoyltransferase I {3 region} [mice, JVS, hearts, Partial, H3140A09678 nt, segment 1 of 2]	major histocompatibility locus class III region:butyrophilin-like protein gene, partial cds; Notch4, PBX2, RAGE, lysophatidic acid acyl transferase-alpha, H3140B07 palmitoyl-protein thioestera	clone RP11-486122, sequence	ectonucleotide pyrophosphatase/phosphodiesterase 5 H3140D05 (putative function) (ENPP5),	H3140D07 peroxiredoxin 3 (Prdx3),	electron-transfer-flavoprotein, alpha polypeptide (glutaric aciduria II) (ETFA),	H3140E04 protein tyrosine phosphatase (70zshp)	glutamate dehydrogenase (Glud),	solute carrier family 12, member 7 (Slc12a7),
H3140A09	H3140B07	H3140C11 clone R	H3140D05	H3140D07	electron polypep H3140D12(ETFA)	H3140E04	H3140E08 glutamat	solute ca H3141F03 (Sic12a7
2787 2788	2789 2790	2791 2792	2793 2794	2795 2796	2797 2798	2799 2800	2801 2802	2803 2804

2805		A - A - on those wortist ofe	Mus musculus chs HMG CoA synthase	BG087678	Mm.10633	Matrix/Structural Proteins
2806	H3141G03 cns HMG "Mus mus	coA syntiase, partia cus sculus protein phosphatase 2a, mbunit, beta isoform (Ppp2cb),				or Thomas de ortion
2808	H3142A02 mRNA"			BG08/094	Mm./410	Digital Maistachou
2809	H3147A08	ubiquinol-cytochrome c reductase core	Homo sapiens uniquinol-cytocia one creductase core protein II (UQCRC2), mRNA	BG075002	Mm.988	Energy/Metabolism
0107						
2811	7010	"Mus musculus tumor-suppressing subchronosomal transferable fragment		BG087715	Mm.139675	Mm.139675 Signal Transduction
2812	H3142E09	1 1 2 2 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1			Mm.31203	Mm.31203 Heat Shock/Stress
		methylenetetrahydrofolate	Homo sapiens methylenetetrahydrofolate dehydrogenase (NADP+ dependent),			
2814		denydrogenase (warta' depondent), methenyltetrahydrofolate cvclohydrolase, myltetrahydrofolate	methenyltetrahydrofolate cyclohydrolase, formyltetrahydrofolate		9	T. Cataladian
2815	H3142G0	H3142G02 synthetase (MTHFD1),	synthetase (MTHFD1), mRNA	BG0877119	Wm.29384	Mm.29384 Energy/Metanousm
2816	H3143D0	H3143D00 heroxisome membrane protein PEX2	Mus musculus peroxisome membrane protein PEX2 mRNA, complete cds	BG075110	Mm.16453	BG075110 Mm.16453 Energy/Metabolism
2818		"Mouse cAMP-dependent protein kinase		,	7	S. Long T. Long of traffice
2819	H3143D1	H3143D10 alpha subunit gene, exon 10"		BG0/2111	MIII.19111	Digual Hansamouron
2820 2821	H3143D1	H3143D12 presenilin-associated protein	Homo sapiens presenilin-associated protein mRNA, complete cds	BG087589	Mm.29650	BG087589 Mm.29650 Energy/Metabolism

2822 2823	H3143H07	"Mus musculus osmotic stress protein H3143H07 94 (Osp94) mRNA, complete cds"		BG075154	Mm.4150	Heat Shock/Stress
2824 2825	H3144B03	NADH-ubiquinone dehydrogenase 1 H3144B03 beta subcomplex	Homo sapiens NADH-ubiquinone dehydrogenase 1 beta subcomplex mRNA, complete cds	BG087636	Mm.4537	Energy/Metabolism
2826 2827	H3144C01	H3144C01 glyoxalase I (GLO1),	ase I (GLO1),	BG087642	Mm.17362	Mm.17362 Energy/Metabolism
2828 2829	"Mus m 5 (gluco H3144C11(Hspa5)	"Mus musculus heat shock 70kD protein 5 (glucose-regulated protein, 78kD) (Hspa5), mRNA"		BG087650	Mm.24162	Mm.24162 Heat Shock/Stress
2830	H3144E06	neuronal cell death related gene in H3144E06 neuron -7 (DN-7)	Rattus norvegicus neuronal cell death related gene in neuron -7 (DN-7) mRNA, complete cds	AW555571 Mm.19440 Apoptosis	Mm.19440	Apoptosis
2831 2832	"Homo s oncogen H3144F09 mRNA"	"Homo sapiens RAB7, member RAS oncogene family-like 1 (RAB7L1), mRNA"		BG075214	Mm.34027	Signal Transduction
2833 2834	H3144G02	"Homo sapiens phosphodiesterase 5A, H3144G02cGMP-specific (PDE5A), mRNA"		BG087768	Mm.103149	Mm.103149 Signal Transduction
2835 2836	H3144H01 mitofilin	mitofilin , partial cds	Homo sapiens mitofilin mRNA, partial cds		Mm.10706	
283 <i>7</i> 2838	H3145A03	"Mus musculus partial Prkar la gene for cAMP-dependent protein kinase regulatory subunit Rialpha, exons 8-10 H3145A03 and 3UTR"		BG075240	Mm.30039	Mm.30039 Signal Transduction
2839 2840	H3145A06	ter ZnT-3 (ZnT-3)	Mus musculus zinc transporter ZnT-3 (ZnT-3) mRNA, complete cds	BG075243	Mm.1396	Energy/Metabolism

		no improve and improve	H samiens wene for phosphate carrier	BG075286	Mm.5246	Energy/Metabolism
	H3145E01 H.sapiens	gene pnospnate carrel	á,	BG075294	Mm.4834	Energy/Metabolism
	113145E11	nyana onemo - ()/)				Heat Shock/Stress
	H3145F11			BG075319	Mm.8385	Signal Transduction
	H3145H03 activated	protein kinase (erk-1)	Mus musculus peroxiredoxin 4 (Prdx4), mRNA	BG075324	_	
2850 2851 2850	H3145H10	H3145H10 neroxitedoxin 4 (Prdx4),	Mus musculus peroxiredoxin 4 (Prdx4), mRNA	BG087872	Mm.19127	Mm.19127 Energy/Metabolism
2853	H3146A04	"Mus musculus myo-inositol 1-phosphate synthase A1 (IsynA1)		BG075331	Mm.29357	Signal Transduction
1007						
2855	102146B10	cDNA FLJ13847 fis, clone THYRO1000852, highly similar to branched-chain amino acid	Homo sapiens cDNA FLJ13847 fis, clone THYRO1000852, highly similar to Human branched-chain amino acid aminotransferase (ECA40) mRNA	BG087895	Mm.24210	Mm.24210 Energy/Metabolism
2857	113146501	lti-drug-	Homo sapiens RHOA proto-oncogene multi-drug-resistance protein mRNA, 3 end	BG087931	Mm.757	Energy/Metabolism
2859	H3146G0t	cDNA: FLJ22871 fis, clone KAT02533	Homo sapiens cDNA: FLJ22871 fis, clone KAT02533		Mm.30065	Energy/Metabolism
2861 2862	H3146H0	H3146H06 CTL2 gene	Homo sapiens CTL2 gene	BG087957	Mm.28209	Energy/Metabolism
2863 2864	H3147B0	"Mus musculus RAB11B, member RAS H3147B08 oncogene family (Rab11b), mRNA"		BG087979	Mm.35727	Mm.35727 Signal Transduction

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147D12		H3147D12 apoptosis inhibitory protein 5 (Api5),	Mus musculus apoptosis inhibitory protein 5 (Api5), mRNA	BG088003 Mm.692	Mm.692	Apoptosis
03271		"Mus musculus signal transducer and activator of transcription 3 interacting		90000000	00030 -31	;; ; ; ; ; ; ; ; ; ; ; ; ; ; ; ; ; ; ; ;
147EC	31 4	∡l ≝IZ			Mm.2180	Digual Hansunction Heat Shock/Stress
147E	5	"Mus musculus heat shock protein, 84 H3147E05 kDa 1 (Hsp84-1), mRNA"		BG088008	Mm.2180	Heat Shock/Stress
147E(9("Mus musculus heat shock protein, 84 H3147E06 kDa 1 (Hsp84-1), mRNA"		BG088009 Mm.2180		Heat Shock/Stress
[48B	8	H3148B04 ferredoxin reductase (Fdxr),	Mus musculus ferredoxin reductase (Fdxr), mRNA	BG088056	4	Energy/Metabolism
(48E	80	H3148E08 ornithine decarboxylase antizyme gene	Mus musculus ornithine decarboxylase antizyme gene, complete eds	BG088091	Mm.683	Energy/Metabolism
H3148F05		NADH dehydrogenase (ubiquinone) Fe- (S protein 1 (75kD) (NADH-coenzyme (Q reductase) (NDUFS1),	Homo sapiens NADH dehydrogenase (ubiquinone) Fe-S protein 1 (75kD) (NADH-coenzyme Q reductase) (NDUFS1), mRNA	BG088100	Mm.2041	Energy/Metabolism
"Mu for H3148G06 cds"	90	"Mus musculus MgcRacGAP mRNA for GTPase activating protein, complete cds"		BG088110	Mm.27141	BG088110 Mm.27141 Signal Transduction
49B	9	"Mus musculus protein kinase C, zeta H3149B10 (Pkcz), mRNA"		BG088144	Mm.28561	BG088144 Mm.28561 Signal Transduction

	1007101	otein, palmitoylated (55	Mus musculus membrane protein, palmitoylated (55 kDa) (Mpp1), mRNA		Mm.2814	Matrix/Structural Proteins
2886	H3149511 H3149C10	H3149C10 (riosephosphate isomerase (Tpi),		BG075608 h	Mm.4222	Energy/Metabolism
2889	H3149E11	уре П,	annel, voltage- eptide	BG088175		Energy/Metabolism
2891	H3149H04	12),	Mus musculus 5-3 exoribonuclease 2 (Xm2), mRNA	BG088199 Mm.3065		Energy/Metabolism
2893	TT2150C0	"Mus musculus Ste-20 related kinase		BG088221	Mm.29404	Signal Transduction
2895 2895 2896	H3150D0	"Mus musculus presentlin 1 (Psen1), H3150D02/mRNA"		BG088227	Mm.998	Signal Transduction
2897	fames;	ransferase, CAAX box, alpha	Mus musculus farnesyltransferase, CAAX box, alpha (Fnta), mRNA	BG075707 Mm.3496	Мт.3496	Energy/Metabolism
2899	H3150G0	: FLJ21905 fis, clone HEP03764	fis,	BG088257	Mm.4290	Energy/Metabolism
2901	H3151C0	"Homo sapiens PTPL1-associated		BG088297		Signal Transduction
2903	H3151D0	H3151D08 defender against cell death 1 (Dad1),	Mus musculus defender against cell death 1 (Dad1), mRNA	BG075765	Mm.2547	Apoptosis
2905	H3151D1		Mus musculus prosaposin (psap\SGP-1) gene, complete cds	BG088310	Mm.3363	Energy/Metabolism
2907 2908	H3151E1	"Mus musculus downstream of tyrosine H3151E12 kinase 1 (Dok1), mRNA"		BG088318	Mm.156	Signal Transduction

2909	H3151F06	H3151F06 M.musculus xanthine dehydrogenase	M.musculus mRNA for xanthine dehydrogenase	BG075778	Mm.11223	Mm.11223 Energy/Metabolism
2911 2912	H3152A02	Rat beta-galactoside-alpha 2,6- H3152A02 sialyltransferase	Rat beta-galactoside-alpha 2,6- sialyltransferase mRNA	BG075800 Mm.1944	Mm.1944	Energy/Metabolism
2913 2914	H3152D06 5-aminol	5-aminolevulinic acid synthase, 3 end	Mouse 5-aminolevulinic acid synthase mRNA, 3 end	BG088378 Mm.1217	Mm.1217	Energy/Metabolism
2915 2916	H3152G07	MAD2 (mitotic arrest deficient, yeast, H3152G07homolog)-like 2 (MAD2L2),	Homo sapiens MAD2 (mitotic arrest deficient, yeast, homolog)-like 2 (MAD2L2), mRNA	:	Mm.9648	Cell Cycle
2917 2918	H3152H05 ATPase,	ATPase, class II, type 9A (Atp9a),	Mus musculus ATPase, class II, type 9A (Atp9a), mRNA	BG075876	Mm.10288	Energy/Metabolism
2918 2919	NADH d beta subc H3152H07 (NDUFB	ehydrogenase (ubiquinone) 1 omplex, 5 (16kD, SGDH) 5),	Homo sapiens NADH dehydrogenase (ubiquinone) 1 beta subcomplex, 5 (16kD, SGDH) (NDUFB5), mRNA	BG088418	Mm.28058	BG088418 Mm.28058 Energy/Metabolism
2920 2921	H3153A02	"Mus musculus calpain small subunit H3153A02[gene, exons 3 through 9 and partial cds"		BG088424 Mm.6534	Mm.6534	Signal Transduction
2922 2923	H3153A07	H3153A07 "Homo sapiens zinedin (ZIN), mRNA"		BG088429	Mm.21612	Signal Transduction
2924 2925	H3153B11 Mm.666	Mm.666			Mm.666	Energy/Metabolism
2926 2927	H3153C09	Mus musculus tissue inhibitor of H3153C09[metalloproteinase 2 (Timp2), mRNA	,	BG088451	Mm.19191	Matrix/Structural Proteins
2928 2930	"Homo s (formerly H3153C10 (65), beta	"Homo sapiens protein phosphatase 2 (formerly 2A), regulatory subunit A (PR 65), beta isoform (PPP2R1B), mRNA"		BG088452	Мт.21428	Mm.21428 Signal Transduction
2931 2932	H3153D05	"Rattus norvegicus Diacylglycerol H3153D05 kinase 90kDa (Dagk), mRNA"		BG075920	Mm.102207	BG075920 Mm.102207 Signal Transduction

2933 2934	p75NT H3153D07 (Nade)	R-associated cell death executor	Mus musculus p75NTR-associated cell death executor (Nade) mRNA, complete cds	BG088461	Mm.90787 Apoptosis	Apoptosis
	NADH d beta subc H3153E04 (NDUFB	l dehydrogenase (ubiquinone) 1 bcomplex, 7 (18kD, B18) FB7),	Homo sapiens NADH dehydrogenase (ubiquinone) 1 beta subcomplex, 7 (18kD, B18) (NDUFB7), mRNA	BG088469	Mm.29683	Energy/Metabolism
2937 2938	M. musc H3153E08 activated	M. musculus mRNA for MAP kinase- activated protein kinase 2		BG088472	Mm.29725	Signal Transduction
2939	apoptotic H3153F08 PtdSerR	cell clearance receptor	Mus musculus apoptotic cell clearance receptor PtdSerR mRNA, complete cds	BG088494	Mm.38825 Apoptosis	Apoptosis
2941 2942	H3153G02 endonuc	lease G-like 1 (ENDOGL1),	Homo sapiens endonuclease G-like 1 (ENDOGL1), mRNA	BG088499	Mm.11669	BG088499 Mm.11669 Energy/Metabolism
2943 2944	"Mus m H3153G08 botulinu	"Mus musculus RAS-related C3 sbotulinum substrate 2 (Rac2), mRNA"		BG088482	Мт.1972	Signal Transduction
2945	U3153H05		Mus musculus peroxisomal delta3, delta2-enoyl-Coenzyme A isomerase (Peci-pending), mRNA	BG075962	Mm.28883	Mm.28883 Energy/Metabolism
2947 2947	H3154A03			BG088511	Mm.7521	Signal Transduction
2949 2950	H3154C11 M.musc	netase	M.musculus mRNA for glutamyl-tRNA synthetase	BG088535	Mm.27190	Mm.27190 Energy/Metabolism
2951 2952	"Mus m H3154E07 (Gas6),	"Mus musculus growth arrest specific 6 (Gas6), mRNA"		BG088548	Мт.3982	Signal Transduction
2953 2954	H3154F02	ine synthetase	Mus musculus asparagine synthetase mRNA, complete cds	BG088553	Mm.2942	Energy/Metabolism

2955 2956	H3155B01	beclin 1 (coiled-coil, myosin-like H3155B01 BCL2-interacting protein) (Becn1),	Mus musculus beclin 1 (coiled-coil, myosin-like BCL2-interacting protein) (Becn1), mRNA	BG088587	BG088587 Mm.30040 Apoptosis	Apoptosis
2957 2958	H3155B07	Homo sapiens NADH dehydrogena: NADH dehydrogena: NADH dehydrogenase (ubiquinone) Fe- (ubiquinone) Fe-S protein 3 (30kD) (NADH-coenzyme (NADH-coenzyme Q reductase) (NDUFS3), (NDUFS3), mRNA	Homo sapiens NADH dehydrogenase (ubiquinone) Fe-S protein 3 (30kD) (NADH-coenzyme Q reductase) (NDUFS3), mRNA	BG076060	Mm.30113	Energy/Metabolism
2959 2960	H3155C04	"Rattus norvegicus mRNA for inositol H3155C04 hexakisphosphate kinase, complete cds"	•	BG076067	Mm.22360	Mm.22360 Signal Transduction
2961 2962	UD H3155C10 06)	P glucuronosyltransferase (UGT1-	Mus musculus UDP glucuronosyltransferase (UGT1-06) mRNA, complete cds	BG076072	Mm.42472	Mm.42472 Energy/Metabolism
2963 2964	H3155E04	H3155E04 NDUFV1 gene, exons 6-10	Homo sapiens NDUFV1 gene, exons 6-10	BG076088	BG076088 Mm.29842	Energy/Metabolism
2965 2966	H3155F02					Heat Shock/Stress
2967 2968	H3155G03	Rat endoplasmic reticulum alpha- H3155G03 mannosidase	Rat endoplasmic reticulum alpha- mannosidase mRNA, complete eds	BG088635	Мт.33138	Mm.33138 Energy/Metabolism
2969 2970	H3156A10					Apoptosis
2971 2972	"Mus mu magnesii H3156C03 (Ppm1d)	"Mus musculus protein phosphatase 1D magnesium-dependent, delta isoform (Ppm1d), mRNA"		BG088678	Мт.61848	Mm.61848 Signal Transduction
2973 2974	H3156C05	CDNA DKFZp586F2224 (from clone H3156C05 DKFZp586F2224)	Homo sapiens mRNA; cDNA DKFZp586F2224 (from clone DKFZp586F2224)	BG076153	Mm.28030	Mm.28030 Energy/Metabolism
2975 2976	H3156C08	H3156C08 "Mus musculus metaxin (Mtx), mRNA"		BG076156	Mm.22508	BG076156 Mm.22508 Heat Shock/Stress

56H04I	Н3156Н04 Н3156Н04	V.			Energy/Metabolism
 ട്ര-	H3157B03 catalase 1 (Cas1),	Mus musculus puranve prospuatase (Pps), mRNA	BG076224 Mm.1458		Energy/Metabolism
- 	"Mus musculus heat shock protein, H3157B10 DNAJ-like 2 (Hsj2), mRNA"		BG088755	Mm.27897	Mm.27897 Heat Shock/Stress
9	(Casp6),	Mus musculus catalase 1 (Cas1), mRNA BG076235		Mm.4215	Energy/Metabolism
8	M.musculus aspartate aminotransferase Mus musculus caspase 6 (Casp6), mRNA mRNA		BG076240 Mm.28814 Apoptosis	Mm.28814	Apoptosis
100	M.musculus aspartate aminotransferase M.musculus aspartate aminotransferase gene exon 10 and 3-flank	M.musculus aspartate aminotransferase gene exon 10 and 3-flank	BG076244	Matrix/S Mm.18916 Proteins	Matrix/Structural Proteins
012	-transferase 3	Homo sapiens microsomal glutathione S-transferase 3 (MGST3), mRNA	BG088778	Мт.29823	BG088778 Mm.29823 Energy/Metabolism
F04	"Mus musculus phosphatidylinositol 3-kinase, catalytic, alpha polypeptide		BG088781	Mm.41943	Mm.41943 Signal Transduction
E05	solute carrier family 30 (zinc H3157E05 (transporter), member 4 (Slc30a4),	Mus musculus solute carrier family 30 (zinc transporter), member 4 (Slc30a4), mRNA	BG076257	Mm.27801	Mm.27801 Energy/Metabolism
H01	H3157H01 sterol O-acyltransferase 1 (Soat1),	Mus musculus sterol O-acyltransferase 1 (Soat1), mRNA	BG088808	Mm.28099	Matrix/Structural Proteins
B04	ornithine decarboxylase antizyme H3158B04 imhibitor (Oazi),	Mus musculus ornithine decarboxylase antizyme inhibitor (Oazi), mRNA	BG088828 Mm.6775		Energy/Metabolism

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10	8C04	Mus musculus maternal embryonic H3158C04 message 3 (Mem3), mRNA		BG076315	Mm 1170	Matrix/Structural
	H3158C07			Cicological		Troteins Heat Chant/Chan
						TICAL DILOCAL DILCOS
	SE03	methylenetetrahydrofolate dehydrogenase (NAD+ dependent), methenyltetrahydrofolate H3158E03 cyclohydrolase (Mthfd2).	Mus musculus methylenetetrahydrofolate dehydrogenase (NAD+ dependent), methenyltetrahydrofolate cyclohydrolase (Ad+442), mDNA	,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	Ş	
	8F09	ne KAT05624	Homo sapiens cDNA: FLJ22903 fis, clone KAT05624	BG088873	Mm 20841	Energy/Metabolism
	8F10	"Mus musculus Ccth gene for chaperonin containing TCP-1 eta H3158F10 subunit, complete cds"		BG076350	Mm.28037	BG076350 Mm.28037 Heat Shock/Stress
~ 1	8G12t	"Mus musculus metallothionein-like 5, H3158G12 testis-specific (tesmin) (Md5), mRNA"		BG076362	Mm 56287	Mm 56287 Heat Shoot/Steem
) H01("Mus musculus protein phosphatase 5 H3158H01(PP5) mRNA, complete cds"				SCALIC DATE OF THE STATE OF THE
	9A03	H3159A03 Mm.10331		1	Mm.3294 Signal Tra	Signal Transduction
	C05 "	"Mus musculus Rab3D (rab3d) gene, H3159C05 complete cds"		BG076387	9 8900 mM	Mm.19951 Apoptosis
\sim 1	D056	"Homo sapiens phosphorylase kinase, H3159D05 beta (PHKB), mRNA"		BG088924	Vim. 28827	BG088924 Mm.28827 Signal Transduction
				1 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2		יופחמן דומווסחחפווחדו

"Mus musculus phosphofructokinase-1 A isozyme (Pfka) mRNA, complete cds" BG088948 Mm.26550 Signal Transduction	Mus musculus glutathione S-transferase, BG088952 Mm.14601 Energy/Metabolism H3159G05 glutathione S-transferase, mu 2 (Gstm2), mu 2 (Gstm2), mRNA		NM 008044 Energy/Metabolism	NM 013671 Energy/Metabolism	NA-12s rRNA	INA-16s TRNA	NA-ATPase 6	NA-ATPase 8	NA-COX 1	NA-COX 2	NA-COX 3	NA-Cyt b	NA-NADH 1	NA-NADH 2	NA-NADH 3	NA-NADH 4	JNA-NADH 4L		NA-NADH 5
Mus musculus phosphofructo , isozyme (Pfka) mRNA, com ds"	lutathione S-transferase, mu	Ant 1	rataxin	· ~	mtDNA-12s rRNA	mtDNA-16s rRNA	mtDNA-ATPase 6	mtDNA-ATPase 8	mtDNA-COX 1	mtDNA-COX 2	mtDNA-COX 3	mtDNA-Cyt b	mtDNA-NADH 1	mtDNA-NADH 2	mtDNA-NADH 3	mtDNA-NADH 4	mtDNA-NADH 4L	mtDNA-NADH 5	
"Mus musc A isozyme H3159F12 cds"	H3159G05g	P1G4	mFRDA Frataxin	CHO3	- 1						1		P1H2		P1H4	PIHS			PIH/
3019 3020	3021 3022		3024		3026	3027	3028	3029	3030	3031	3032	3033	3034	3035	3036	3037	3038		3039

Table 5

Num		Genbank Acc. #	IMAGE Clone ID	Unigene Cluster	Res. Gen.Clone
	mtDNA-12s rRNA	V00711	In House-Clone		
	mtDNA-16s rRNA	V00711	In House-Clone		
	mtDNA-ATPase 6	V00711	In House-Clone		
	mtDNA-ATPase 8	V00711	In House-Clone		
	mtDNA-COX 1	V00711	In House-Clone		
	mtDNA-COX 2	V00711	In House-Clone		
	mtDNA-COX 3	V00711	In House-Clone	 	
	mtDNA-Cyt b	V00711	In House-Clone		
	mtDNA-NADH 1	V00711	In House-Clone		
	mtDNA-NADH 2	V00711	In House-Clone		
	mtDNA-NADH 3	V00711	In House-Clone		
	mtDNA-NADH 4	V00711	In House-Clone		
	mtDNA-NADH 4L	V00711	In House-Clone		
	mtDNA-NADH 5	V00711	In House-Clone		
	mtDNA-NADH 6	V00711	In House-Clone		
13	INDIVITUAL O	X04958,	1110000 010110	- ·	
16	(2'-5')oligoadenylate synthetase 1	M33863	AI449562	Mm.14301	577664
	14-3-3 protein beta	AF058797.1	AA624796	Mm.34454	1039350
	14-3-3 protein epsilon	D87663.1	AA240827	Mm.678	670982
	14-3-3 protein eta	U57311	AI325059	Mm.3308	558536
	14-3-3 protein gamma	AF058799	122200	Mm.10802	464892
	14-3-3 protein theta (tau)	U57312	AA422980	Mm.14722	803843
	14-3-3 protein zeta	U79231	AA671451	Mm.3360	1039129
23	25-hydroxyvitamin D3 24-hydroxylase precursor	D49438	AA240836	Mm.6575	657172
24	25-hydroxyvitamin D3 24-hydroxylase precursor	D49438	AI226268	Mm.6575	658678
25	2-amino-3ketobutyrate-coenzyme A ligase	AF093403	AI037094	Mm.18618	315143
25	2-oxoglutarate dehydrogenase E1	1700071	77/00/442		10.1.150
26	component	U02971	W98443		424460
27	3,2-trans-enoyl-CoA isom, mito precursor	Z14049	AI194961		1886651
28	38g cent.		In House-Clone		
29	3-beta hydroxy-5-ene steroid dehydrogenase type I	M27137	AA274685		766591
30	3-beta hydroxy-5-ene steroid dehydrogenase type II	M75886	AI266804	Mm.30433	1891212
31	<u> </u>	M77015	AA209060	Mm.335	676577
32		L16992		Mm.12819	422840
	41.2a cent.		In House-Clone		
34			In House-Clone		ļ
35	45.2b cent.		In House-Clone		
36	55e cent.	L	In House-Clone		
37	5-aminolevulinate synthase precursor (EST)	M15268	AA189529	Mm.1217	635215
38	65c cent.		In House-Clone		

11					
39	66a cent.		In House-Clone		
	67 cent.		In House-Clone		
	a-amylase	V00719	AI325237	Mm.7074	608852
	acatyl-CoA acetyltransferase 1	L42293		Mm.28099	472233
	acatyl-CoA acetyltransferase 2	AF078751	AA239043		694062
44	Aconitase (iron responsive element)	X61147	AA212704		677092
45	Aconitase (iron responsive element)	X61147	AA238899		697949
-10	Acyl-CoA dehyd, med-chain specific				
46	precursor (MCAD)	U07159	AA104184	Mm.10530	568149
	Acyl-CoA dehyd, short-chain specific				
47	precursor	L11163	AI050239	Mm.18759	1379035
	Acyl-CoA dehyd, very-long-chain specific				
48	precursor	U41497	AA250410	Mm.18630	670916
	Acyl-CoA dehydrog, long-chain spec.		,		
49	precursor (LCAD)	U21489	AA254905	Mm.2445	719580
	adapt		In House-Clone		
	Adenine nucleotide translocator 2,				
51	fibroblast (Ant2)		AA033138.1		465520
52	adenylate kinase isoenzyme 2 (EST)	AI155541	AA061587	Mm.29460	483322
53	Adenylosuccinate synthase		AA388461		749837
54	adenylyl cyclase type VII	U12919	W65619	· · · · · · · · · · · · · · · · · · ·	387280
24	ADP, ATP carrier protein, fibroblast				
55	isoform 2 (ant2)	U27316		Mm.658	585992
-33	isotoria 2 (data)				
56	ADP, ATP carrier protein, heart isoform T1	U27315	AA717872	Mm.16228	1152250
57	adrenodoxin precursor	L29123	AA461849	Mm.1061	851558
58	Alcohol dehydrogenase 5		AA183192		636207
59	Alcohol dehydrogenase I		AA221141		695105
60	Aldehyde dehydrogenase	U07235	AA122975		579570
	aldehyde dehydrogenase (NAD+) 2				
61	precursor	U07235	AI503977	Mm.2621	1001020
62	Alpha-1 protease inhibitor	U38477	AA212578	Mm.16672	676745
63	Ant 1		In House-Clone		
64	Antioxidant protein 1	M28723	W91307		423832
65	Antioxidant protein 1	M28723		Mm.29821	599017
66	Antioxidant protein-2 (AO2)		AA243957	Mm6587	694088
67	Apoptosis inhibitor 2		AA144490		597715
68			AA097958		550702
1	apoptosis regulator BAX, membrane				
69	LE E LE	L22472	AI323521	Mm19904	557643
70		AF100927	AA866777		1434491
71	ASC		In House-Clone		
·-	ATP synth lipid-binding protein P1				
72	1 1 1 1 1	L19737	AI481739	Mm.258	888863
73		AA106406	AA106406	Mm.5293	519329
74		L19737	AA139793		580898
75			AA269701		735887
76			AA184876		642977
	ATPase, Ca++ transporting, cardiac				
77		<u> </u>	W34420.1		318735
78			W34420		318735
79			AA276030		776055
80		U43892	AU019072	Mm.4739	1920872

Branched chain alpha ketoacid dehydrog. Mm.8903 578018	81	Bcl-2 protein	U10102	AA051441		478723
branched chain alpha-ketoacid dehyd chain L47335 Mm.25848 314098						
Branched-chain a-ketoacid dehydrog, E1b AA050586 476163					Mm.8903	578018
Branched-chain a-ketoacid dehydrog. E1b					Mm.25848	314098
Barached-chain amino transferase I AA003372 A26976						
Section				AA050586		476163
ST CII 3				AA003372		426976
Section						
Calcium channel, voltage-dependent, L ype, alpha 2 delta subunit In House-Clone						
Supe, alpha 2 delta subunit	88			In House-Clone	<u> </u>	
91 Calmodulin 3 S71789 S2 Carbonate dehydratase, hepatic X51971 A1256540 Mm.35538 1889415 Mm.26141 1167078 A270101 AA71129 Mm.1614 1167078 A270101 AA71129 Mm.21454 317572 A270101 A2701		type, alpha 2 delta subunit		AA511037.1		890932
92				In House-Clone		
93 Carbonic Anhydrase IV U37091 AA71129 Mm.1614 1167078 94 Carbonyl reductase (NADPH) - mouse D26123 W11423 Mm.21454 317572 95 Carnitine O-acetyltransferase (carnitine of acetylase) X85983 AI528757 Mm.20396 602213 96 Carnitine O-palmitoyltransferase II D1163 Mm.29499 580316 97 Carnitine palmitoyltransferase 2 Mm.29499 580316 98 Caspase 2 AA200808 G39403 S50766 100 Catalase AA239490 Mm 4215 678773 CCAAT/enhancer binding protein CCBP), alpha AA271223.1 738252 102 CEBP In House-Clone D1630 Mm.12970 422572 105 CI 18K In House-Clone D1630 Mm.12970 422572 106 CI 18 In House-Clone D1630 Mm.25820 390487 101 Coproporphyrinogen III oxidase precursor D16333 W71884 Mm.35820 390487 102 Coproporphyrinogen oxidase D16333 AA108600 570602 103 Core binding factor beta Carabase D16333 AA259342 734795 114 Core binding factor beta Carabase D16333 AA168601 Mm.970 1167886 115 Creatine kinase Carabase D16333 AA166212 Mm.2375 608246 116 Creatine kinase - complete M74149 AA270310 Mm.16831 736251 117 Creatine kinase - mitochondrial Carabase C				AA109041.1		571789
Searbonyl reductase (NADPH) - mouse D26123 W11423 Mm.21454 317572			X51971	AI256540	Mm.35538	1889415
24	93	Carbonic Anhydrase IV	U37091	AA71129	Mm.1614	1167078
Camitine O-acetyltransferase (carnitine acetylase) X85983 AI528757 Mm.20396 602213	94	carbonyl reductase (NADPH) - mouse		W11423	Mm.21454	317572
Carnitine O-palmitoyltransferase II Frecursor U01163 Mm.29499 S80316		carnitine O-acetyltransferase (carnitine				
96 precursor U01163 Mm.29499 580316 97 Carnitine palmitoyltransferase 2 AA200808 639403 98 Caspase 2 AA200808 639403 99 Caspase I AA098139 550766 100 Catalase AA239490 Mm 4215 678773 CCAAT/enhancer binding protein AA271223.1 738252 101 (C/BBP), alpha AA271223.1 738252 102 CEBP In House-Clone 102 103 Cellular apoptosis succep. Protein AA471761 874148 104 chaperonin-10 U09659 Mm.12970 422572 105 CI 18K In House-Clone In House-Clone 106 CI B8 In House-Clone In House-Clone 109 coproporphyrinogen oxidase D16333 W71884 Mm.35820 390487 110 coproporphyrinogen oxidase D16333 AA108600 570602 111 coproporphyrinogen oxidase D16333 W53951 367358 </td <td>95</td> <td>· · · · · · · · · · · · · · · · · · ·</td> <td>X85983</td> <td>AI528757</td> <td>Mm.20396</td> <td>602213</td>	95	· · · · · · · · · · · · · · · · · · ·	X85983	AI528757	Mm.20396	602213
97 Carnitine palmitoyltransferase 2	96		U01163	1	Mm 29499	580316
98 Caspase I AA200808 639403 99 Caspase I AA098139 550766 100 Catalase AA239490 Mm 4215 678773 CCAAT/enhancer binding protein CCAAT/enhancer binding protein AA271223.1 738252 102 CEBP In House-Clone B74148 103 Cellular apoptosis succep. Protein AA471761 874148 104 chaperonin-10 U09659 Mm.12970 422572 105 CI 18K In House-Clone In House-		h				
99 Caspase I AA098139 550766 100 Catalase AA239490 Mm 4215 678773 CCAAT/enhancer binding protein CCABT/enhancer binding protein AA271223.1 738252 101 (C/EBP), alpha AA471761 874148 102 CEBP In House-Clone 874148 104 chaperonin-10 U09659 Mm.12970 422572 105 CI 18K In House-Clone 106 CI B8 In House-Clone 107 col 1 In House-Clone 107 col 1 108 col 3 109 coproporphyrinogen III oxidase precursor D16333 W71884 Mm.35820 390487 110 coproporphyrinogen oxidase D16333 AA108600 570602 111 coproporphyrinogen oxidase D16333 AA259342 734795 112 coproporphyrinogen oxidase D16333 W53951 367358 113 coproporphyrinogen oxidase D16333 W53951 367358 114 creatine kinase Z13968 AA146442.1 596552 114 creatine kinase, muscle X03233 AA166212 Mm.2375 608246				AA200808		
100 Catalase						
CCAAT/enhancer binding protein (C/EBP), alpha		<u> </u>			Mm 4215	
101 C/EBP), alpha		CCAAT/enhancer binding protein				0,0,7,5
102 CEBP	101			AA271223.1		738252
104 chaperonin-10 U09659 Mm.12970 422572 105 CI 18K In House-Clone 106 CI B8 In House-Clone 107 108 In House-Clone 109 109 100	102	CEBP		In House-Clone		
In House-Clone In House In House In House In House In House In House In House In House In House In House In House In House In House In House In House In House In House In House In House In House In House In House In House In House In House In House In House In House In House In House In House In House In House In House In House In House In House In House In House In House In House In House In House In House In House In House				AA471761		874148
In House-Clone In H			U09659		Mm.12970	422572
In House-Clone In House-Clone 108 col 3 In House-Clone 109 coproporphyrinogen III oxidase precursor D16333 W71884 Mm.35820 390487 110 coproporphyrinogen oxidase D16333 AA108600 570602 111 coproporphyrinogen oxidase D16333 AA259342 734795 112 coproporphyrinogen oxidase D16333 W53951 367358 113 Core binding factor beta AA146442.1 596552 213968 AA146442.1 596552 114 creatine kinase Z13968 AA690010 Mm.970 1167886 115 Creatine Kinase, muscle X03233 AA166212 Mm.2375 608246 116 Creatine kinase-complete M74149 AA270310 Mm.16831 736251 117 Creatine kinase-mitochondrial Z13968 AI528837 Mm.970 607301 cyt C oxydase polypeptide VIIa-liver/heart precursor X58486 AA960158 Mm.2151 1248366 cyto C oxydase polypeptide VIa-heart U08439 AA415934 Mm.21050 846138 cyto C oxydase polypeptide VIa-liver Coxydase polypeptide VII-liver Coxydase polypepti				In House-Clone		
108 col 3						
109		<u> </u>				
110 coproporphyrinogen oxidase D16333 AA108600 570602 111 coproporphyrinogen oxidase D16333 AA259342 734795 112 coproporphyrinogen oxidase D16333 W53951 367358 113 Core binding factor beta AA146442.1 596552 114 creatine kinase Z13968, AA690010 Mm.970 1167886 115 Creatine Kinase, muscle X03233 AA166212 Mm.2375 608246 116 Creatine kinase-complete M74149 AA270310 Mm.16831 736251 117 Creatine kinase-mitochondrial Z13968 AI528837 Mm.970 607301 cyt C oxydase polypeptide VIIa- X58486 AA960158 Mm.2151 1248366 cyto C oxydase polypeptide VIa-heart U08439 AA415934 Mm.21050 846138	108	col 3		In House-Clone		
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112 coproporphyrinogen oxidase D16333 W53951 367358 113 Core binding factor beta AA146442.1 596552 213968, Z13968, AA690010 Mm.970 1167886 115 Creatine kinase X03233 AA166212 Mm.2375 608246 116 Creatine kinase-complete M74149 AA270310 Mm.16831 736251 117 Creatine kinase-mitochondrial Z13968 AI528837 Mm.970 607301 cyt C oxydase polypeptide VIIa- X58486 AA960158 Mm.2151 1248366 cyto C oxydase polypeptide VIa-heart U08439 AA415934 Mm.21050 846138						570602
113 Core binding factor beta AA146442.1 596552 114 creatine kinase Z13968, Z13969 AA690010 Mm.970 1167886 115 Creatine Kinase, muscle X03233 AA166212 Mm.2375 608246 116 Creatine kinase-complete M74149 AA270310 Mm.16831 736251 117 Creatine kinase-mitochondrial Z13968 AI528837 Mm.970 607301 cyt C oxydase polypeptide VIIa- Iliver/heart precursor X58486 AA960158 Mm.2151 1248366 cyto C oxydase polypeptide VIa-heart Ilip precursor U08439 AA415934 Mm.21050 846138 cyto C oxydase polypeptide VIa-liver Coxydase polypeptide VII-liver Coxydase polypeptide VII-live					•	
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114 creatine kinase Z13969 AA690010 Mm.970 1167886 115 Creatine Kinase, muscle X03233 AA166212 Mm.2375 608246 116 Creatine kinase-complete M74149 AA270310 Mm.16831 736251 117 Creatine kinase-mitochondrial Z13968 AI528837 Mm.970 607301 cyt C oxydase polypeptide VIIa-liver X58486 AA960158 Mm.2151 1248366 cyto C oxydase polypeptide VIa-heart U08439 AA415934 Mm.21050 846138 cyto C oxydase polypeptide VIa-liver TAA415934 Mm.21050 846138	113	Core binding factor beta		AA146442.1		596552
115 Creatine Kinase, muscle X03233 AA166212 Mm.2375 608246 116 Creatine kinase-complete M74149 AA270310 Mm.16831 736251 117 Creatine kinase-mitochondrial Z13968 AI528837 Mm.970 607301 cyt C oxydase polypeptide VIIa-liver X58486 AA960158 Mm.2151 1248366 cyto C oxydase polypeptide VIa-heart U08439 AA415934 Mm.21050 846138 cyto C oxydase polypeptide VIa-liver Coto Coto oxydase polypeptide VIa-liver Coto Coto oxydase polypeptide VIa-liver Coto Coto oxydase polypeptide VIa-liver						
116 Creatine kinase-complete M74149 AA270310 Mm.16831 736251 117 Creatine kinase-mitochondrial Z13968 AI528837 Mm.970 607301 cyt C oxydase polypeptide VIIa- liver/heart precursor X58486 AA960158 Mm.2151 1248366 cyto C oxydase polypeptide VIa-heart 119 precursor U08439 AA415934 Mm.21050 846138 cyto C oxydase polypeptide VIa-liver TAA415934 Mm.21050 846138						
117 Creatine kinase-mitochondrial Z13968 AI528837 Mm.970 607301 cyt C oxydase polypeptide VIIa- 118 liver/heart precursor X58486 AA960158 Mm.2151 1248366 cyto C oxydase polypeptide VIa-heart 119 precursor U08439 AA415934 Mm.21050 846138 cyto C oxydase polypeptide VIa-liver		· · · · · · · · · · · · · · · · · · ·		 		
cyt C oxydase polypeptide VIIa- 118 liver/heart precursor X58486 AA960158 Mm.2151 1248366 cyto C oxydase polypeptide VIa-heart 119 precursor U08439 AA415934 Mm.21050 846138 cyto C oxydase polypeptide VIa-liver				· · · · · · · · · · · · · · · · · · ·		
118 liver/heart precursor X58486 AA960158 Mm.2151 1248366 cyto C oxydase polypeptide VIa-heart U08439 AA415934 Mm.21050 846138 cyto C oxydase polypeptide VIa-liver Control oxydase polypeptide VIa-liver Control oxydase polypeptide VIa-liver Control oxydase polypeptide VIa-liver	11/		Z13968	A1528837	Mm.970	607301
119 precursor U08439 AA415934 Mm.21050 846138 cyto C oxydase polypeptide VIa-liver TAMES TO SERVICE TO SERVIC	118	liver/heart precursor	X58486	AA960158	Mm.2151	1248366
cyto C oxydase polypeptide VIa-liver	119		U08439	AA415934	Mm.21050	846138
1001				1		0,0100
	120		L06465		Mm.19094	533628

•					
ì	\	X54691,			
121	cytochrome c oxidase chain IV precursor	M37829	AA260009	Mm.2136	746546
122	cytochrome c oxidase chain Va precursor	X15963	AI131665	Mm.360	1884978
123	cytochrome c oxidase chain Vb precursor	X53157	AI035302	Mm.16769	1432845
	cytochrome c oxidase chain VIb				
	HUMAN)	NM 001863	AI503861		991287
125	cytochrome c oxidase chain VIc	M20153	AA062417		516889
125	cytochrome c oxidase chain VIIc	X52940	AA031250	Mm.14831	464966
127	cytochrome c oxidase chain VIII	U37721		Mm.14022	481408
120	Cytochrome C oxidase sub VII		AA050684		476180
	cytochrome C oxydase polypeptide VIII-H				
	precursor	U15541	AI323348	Mm.3841	463967
	cytochrome C, somatic	X01756	AA221965	Mm.35389	658678
130	cytochrome C-type heme lyase (CCHL)	U36778		Mm.3988	400735
131	cytochrome-b5 reductase (HUMAN)	NM 000398	AA816039		1120651
132	~ <i>_</i>	1111_000350	12101000		
	CYTOSOLIC BRANCH CHAIN AMINOTRANSFERASE		AA286063		776036
			In House-Clone		7,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,
	DD 43		In House-Clone		
	DD 47	 	In House-Clone		
	DD 48		In House-Clone		
	DD 53		In House-Clone		
	DD 64		In House-Clone		
	DD 68	-	In House-Clone	·	
	DD 69		In House-Clone		+
	DD 73	 	In House-Clone		
	DD 83	}	 		-
	DD 84	<u> </u>	In House-Clone		-
	DD10a		In House-Clone		
	DD11b	 	In House-Clone In House-Clone		+
	DD14	 			
	DD15c		In House-Clone		
	DD16c	 	In House-Clone		
	DD17c	- 	In House-Clone		
	DD19a	 	In House-Clone		
	DD23ba		In House-Clone		-
152	DD24d		In House-Clone		<u> </u>
	DD25c	- 	In House-Clone	·	
	DD26f		In House-Clone		
	DD2a	-	In House-Clone		
156	DD33a/pgf		In House-Clone		
157	DD39		In House-Clone		
	DD4c	 	In House-Clone		
159	DD7a		In House-Clone		161600
160	defender against cell death 1		AA033006		464622
161	DiGeorge syndrome chromosome region	6	W54234.1		356181
	dihydrolipoamide dehydrogenase (E3)				00.400
163	(HUMAN)	J03490	AA548170	Hs.74635	994825
	dihydrolipoamide transacylase precursor	L/42996	AA254971	Mm.3636	719973
	Dimethyl glycine dehydrogenase		AA288418		748958
	5 DNA polymerase gamma	U53584		Mm.3616	575332
	6 Dynamin		AA266438	<u> </u>	317587

167	Dynamin 2		W13111		457445
	E1B 19k/Bcls-binding prot. Homolog				İ
	(NIP3)		AA105295		571367
	ER V		In House-Clone		
	ERV-1	U40494	AI413376	Mm.28124	367232
	EST highly similar to S-100 prot a-chain		AA466432		872869
	ESTs		AA253853.1		660997
	ESTs		AA268402.1		721970
174	ESTs		W14142.1		329863
175	ESTs, Highly similar to CREATINE KINASE, SARCOMERIC MITO. PRECUR.		AA038095.1		472860
176	ESTs, Highly similar to CREATINE KINASE, SARCOMERIC MITO. PRECUR.		AI322288.1		336085
177	ESTs, Highly similar to DERMATOPONTIN [Bos taurus]		W13931.1		330218
178	ESTs, Highly similar to MATERNAL EFFECT PROTEIN STAUFEN		AA104976.1		533314
179	ESTs, Highly similar to NADH- UBIQUINONE OXIDOREDUCTASE 13 KD-B SUB		AA288040.1		748891
180	ESTs, Highly similar to NADH- UBIQUINONE OXIDOREDUCTASE SGDH SUB PREC.		AA222463.1	•	671212
181	ESTs, Highly similar to NUCLEAR FACTOR NF-KAPPA-B P100 SUBUNIT		AA060802.1		482952
182	ESTs, Highly similar to PINCH PROTEIN		AA289280.1		790449
183	ESTs, Highly similar to PTERIN-4- ALPHA-CARBINOLAMINE DEHYDRATASE		W14332.1		331681
184	ESTs, Highly similar to PUTATIVE REGULATORY PROTEIN TSC-22		W66757.1		387449
185	ESTs, Highly similar to SUCCINATE DEHYDROGENASE		AA108475.1		572339
186	ESTs, Highly similar to (defline not available 4588044)		AA190123.1		642467
187	ESTs, Highly similar to putative ATP/GTP-binding protein		AA184876.1		642977
188	ESTs, Weakly similar to (defline not available 3668141) [H.sapiens]	•	AA467585.1	•	833160
189	ESTs, Weakly similar to survival motor neuron [M.musculus]		AA538419.1		932748
	ESTs, Weakly similar to vesicle membrane protein		AA259674.1		735186
	Excitatory amino acid transporter 3	D43797	AA065502	Mm.24741	524046
	ferredoxinNADP+ reductase precursor	D49920	AA879949	Mm.4719	1230740
	ferrochelatase precursor	M59288		Mm.1070	635652
	Friedreich ataxia	S75712	AA930748		1150363
	Fructose 1-6 bis-phosphate	D42083	AI385602	Mm.2974	336727
	Fructose bis-phosphate aldolase	W53351	AI553136	Mm.7729	902910
197	Fructose Bisphosphate aldolase A	Y00516	AA518639		903419

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198	Fuzzy onion homolog (mouse)		AA674474		1093002
199	GAMMA-ADAPTIN		AA238435.1		693837
200	GAPD		In House-Clone		
201	GILZ		In House-Clone		
	Gluamate receptor, ionotropic, kainate 5 (gamma 2)		AA261334	Mm.2879	733368
203	Gluamate receptor, ionotropic, NMDA1 (zeta 1)		W44130	Mm.3292	354244
204	Gluamate receptor, ionotropic, NMDA2C epsilon 3)		AI256808	Mm.39090	1852361
	Glucose dependent insulinotropic polypeptide	U34295	AA871367	Mm.5115	1096156
206	Glucose Phospahate Isomerase I complex	M14220	AA276216		776210
207	Glucose Phospahate Isomerase I complex	M14220	W29397		337413
208	Glucose-6-phosphate isomerase	L09104	AI327180	Mm.589	437357
	glutamate dehydrogenase (NAD(P)+) precursor	X57024	AA543797	Mm.10600	949005
210	glutamate oxaloacetate transaminase-2	X06917, J02622		Mm.18916	617490
	Glutaryl-CoA dehydrogenase precursor (GCD)	U18992		Mm.2475	573351
212	Glutathione peroxidase-3 (plasma Gpx)	U13705	AA097557	Mm7156	552393
213	Glutathione peroxidase-4 (phospholipid Gpx)		AA034666	Mm2400	466963
214	Glutathione peroxidase-heart isoform (Gpx4)	AF045769.1	AI327053	Mm.2400	420345
215	Glutathione peroxidase-plasma isoform	U13705.1	AI042912	Mm.7156	1432410
216	Glyceraldehyde 3-phosphate dehydrogenase		AA122891		579715
217	Glycerol 3-phosphate acyltransferase	M77003	AA209041		676437
	Glycerophosphate dehydrogenase	M13366	AI414023	Mm.10669	303389
	Glycogen Phosphorylase (RAT)	J03080	AA240684		656882
	GP4		In House-Clone		
	Gpx		In House-Clone		
	GTP2		In House-Clone		
223	GTP-binding protein		AA509565		890444
224	H+-transporting ATP synthase chain alpha	L01062	AI573940	Mm.4069	1920315
225	H+-transporting ATP synthase chain beta	AF030559	AI452208	Mm.17869	576006
226	H+-transporting ATP synthase chain delta (EST)	AA940343	AA896697	Mm.22514	1279053
	H+-transporting ATP synthase chain e	U59283	AI115240		1885128
	H+-transporting ATP synthase chain e	U59283	AI481625		876416
	H+-transporting ATP synthase chain gamma	U43893	AA210528		640931
	heat shock protein 60 precursor	X55023, X53584	AA154729	Mm.1777	540881
	heat shock protein 70 precursor	D11089	AI132204	Mm.2849	1481709
	heat shock protein, 70K (hsp68) (fragment)	M12573	AA647374	Mm.6388	1108306
	Heat shock protein, 84 kDa 1	M18186	141011317	Mm.2180	538585
	Hexokinase	1	W11571		318642

235	Hexokinase	1	W74835	1	389177
236	hexokinase I (3'-seq)	J05277	AI661880	Mm.5290	717383
237	hexokinase I (5'-seq)	J05277	AA197916	Mm.5290	654873
	high mobility group protein homolog				+ 30.075
238	HMG4 (Hmg4) mRNA, complete cds		AA048831.1		478865
239	HK-40s Ribosomal protein S15		AA033398		466295
240	HK-40s Ribosomal protein S4		AA000082		425352
241	HK-A111Acient ubiquinating protein		W82989		405768
242	HK-A16060s Ribosomal protein L15		AA068842		532770
243	HK-A17360s Ribosomal protein L3		AA108363		570533
244	HK-A21060s Ribosomal protein L1A		AA170607		619006
	HK-A216RNA splicing protein		AA183061		636766
	HK-A262E2F transcription factor		AA396123		751755
	HK-A272Capping protein		AA414612		779754
	HK-A316Actin-gamma (smooth muscle)	M26689	AA710883	Mm.16562	1166855
	HK-A97Hypozanthine phosphoribosyl		12171000	171111.10502	1100855
	transferase	J00423	W48168	Mm.18675	355084
250	HK-Actin- alpha (skeletal muscle)	M12347	AI035279	Mm.4581	1480709
	HK-Actin- gamma	M21495	AI314957	Mm.29913	192859
	HK-Alkaline Phosphatase		AA032457	112111255715	465052
	HK-b-actin	X03672	AA079937	Mm.297	536615
	HK-calcium binding protein Cab45	U45977	AI266799	Mm.30149	1891202
	HK-DNA ligase I		W66626	MILOUT	388245
	HK-Glyceraldehyde 3-phosphate		W 00020		366243
	dehydrogenase (G3PDH)	W14827	AA119563	Mm.5289	538210
	HK-Glyceraldehyde phosphate		12222000	14111.0209	338210
	dehydrogenase	M32599	AA466618	Mm.5289	817984
258	HK-HPRT		In House-Clone		02.701
	HK-Hypozanthine phosphoribosyl				
259	transferase	J00423	AI256193	Mm.18675	1890233
260	HK-Murine ornithine decarboxylase	M10624	AI325192	Mm.15259	608003
261	HK-Myosin 1	L00923	AI098184	Mm.3390	1481939
	HK-Phospholipase A2 (14-3-3 zeta/delta)	D78647	AA1714341	Mm.3360	617315
	HK-Ribosomal Protein S29	L31609	AA032465	Mm.35816	465138
264	HK-Ubiquitin	X51703	AI181949	Mm.235	1451597
	House mouse; Musculus domesticus				
	mRNA for LN1, complete cds		W82212.1		403728
	Huntingtin	AI876894	AI876894		1922060
	hydroxymethylglutaryl-CoA lyase	S65036	AA838929		1261134
	hydroxymethylglutaryl-CoA synthase,	U12790,			
	mitochondrial	U12791		Mm.10633	518481
	IG ALPHA CHAIN C REGION		AA098196.1		551003
	IMAGE EST		AA009059.1		441176
	IMAGE EST		AA028306		464099
	IMAGE EST		AA035899.1		468817
	IMAGE EST		AA051664.1		479709
-	IMAGE EST		AA118290		574435
_	IMAGE EST		AA200984.1		639212
	IMAGE EST		AA203878		640734
-	IMAGE EST		AA215024		652207
	IMAGE EST		AA221250		670393
279	IMAGE EST		AA245545		699280

•				1	
280 lt	MAGE EST		AA250652		697537
	MAGE EST		AA266097.1		716941
	MAGE EST		AA275684		775722
	MAGE EST		AA388512		790857
	MAGE EST		AA466026.1		809016
	MAGE EST		AA519027.1		904900
	MAGE EST		W08090.1		331768
	MAGE EST		W09924		315773
	MAGE EST		W09924.1		315773
	MAGE EST		W15031		330502
	MAGE EST		W41309.1		351420
	IMAGE EST		W89418.1		420553
291	IMAGE EST- CCAAT enhancer binding				
	protein		AA271223		738252
292	IMAGE EST-Glucocorticoid-induced				
202	leucine zipper GILZ protein	ł	W66757		387449
	IMAGE EST-homolog of Unc33 (C.Elegans) / Collaspin reponse mediated				
204	prot. 2		W08090		331768
294	IMAGE EST-sarcoplasmic creatine kinase		W18057		336085
293	IMAGE EST-Sim to gamma sarcoglycan		W41309		351420
296	IMAGE EST-yeast bile transporter		AA473289		803488
	Interferon gamma receptor		AA541842.1		920516
298	Interleukin 1 receptor-associated kinase		AA276835.1		777580
299	isocitrate dehydrogenase (NADP)	U51167		Mm.2966	571468
			In House-Clone		
	Kin 17	M27554	AA880398		1277670
302	Lactate Dehydrogenase- Lactate Dehydrogenase-A4	M17516	AI506641	Mm.26504	1024774
303	Lactate Denydrogenase-A-	M17587,			
	Lactate Dehydrogenase-sperm specific	L10389	AA110449	Mm.16563	516582
304	M.musculus mRNA for fibromodulin		W63981.1		374228
303	M.musculus mRNA for GTP-binding				
206	protein protein	1	AA020462.1		455401
300	MAD homolog 4 (Drosophila)		AA030901.1		466551
307	Malate dehydrogenase (cyto)		W13686		318346
	Malate dehydrogenase (mito)	M16229	AA266087	_	717095
309	malate dehydrogenase precursor,	X07295,			
210	mitochondrial	M16229		Mm.21743	407143
211	MAP KINASE PHOSPHATASE-1		AA125367.1		575665
311	Maternal embryonic message 3		AA388122.1		775464
	MCK.		In House-Clone		
	mDP 6	<u> </u>	In House-Clone		
314	metal response element DNA-binding	1			
31	protein M96 mRNA, complete cds	1	AA545607.1		945218
313	methylenetetrahydrofolate dehydrogenase				
21	methylenetetranydrofolate denydrogenase 5 (NAD+)	J04627		Mm.443	440345
31	methylmalonyl-CoA mutase alpha chain	T			
1	methylmalonyi-CoA mutase alpha cham precursor	X51941	1	Mm.4299	571282
31	7 precursor 8 Microtubule-associated protein 4		AA003769.1		437523
31	9 Mito matrix prot P1 precursor (hsp60)	1	AA184322		633625
31	O Miles handrial I ON protesse	AA061310	AA061310		514859
32	0 Mitochondrial LON protease	+	1		
	mitochondrial transcription factor A	U57939		Mm.276	539693
32	1 mouse	U57939	1	IVIII.270	1 33909

322	mitochondrial uncoupling protein	M21247	AI131780	Mm.4177	1498957
	monoamine oxidase A - mouse	S78615	AI643185		864614
324	Mouse breast heat shock 73 prot (hsc73)	M19141		Mm.2944	538418
	Mouse calcineurin catalytic subunit				
325	mRNA, complete cds		AA245461.1		699236
	Mouse Circadian output locomotor cycles				
	kaput	AF000998	AI156715	Mm.3552	1494023
	Mouse heatshock protein 27	U03560.1	AA596241	Mm.13849	1052188
-	Mouse heatshock protein 86	J04633	AI649095		1970053
	Mouse med. Chain acyl-CoA	7705150	11061670		
	dehydrogenase	U07159	AA061679	3.6 40.60	483333
	Mouse mHox protein	L06502	W17990	Mm.3869	335936
331	Mouse Skd3 mRNA	U09874		Mm.3990	602340
	mRNA for sarco/endoplasmic reticulum Ca2+ -ATPase (SERCA2)	·	AA222567.1		695695
333	mTF 1		In House-Clone		
	Mus musculus Balb/c zinc finger protein				
334	PZF (Pzf) mRNA, complete cds		W11161.1		316427
335	Mus musculus calcium-binding protein S100A1 mRNA, complete cds		AA466432.1		872869
	Mus musculus cytoplasmic protein Ndr1				0,200
336	(Ndr1) mRNA, complete cds		AA473269.1		803416
337	Mus musculus FGF-binding protein (FGF-BP) mRNA, complete cds		AA403432.1		717457
	Mus musculus GTP binding protein (GTP2) mRNA, complete cds		AA509565.1		890444
338	(G112) midvA, complete cus		AA309303.1		090444
	Mus musculus hemin-sensitive initiation				•
339	factor 2 alpha kinase mRNA, complete cds		AA036546.1		466971
	Mus Musculus mRNA for glutamate				
340	receptor channel alpha 4 subunit		AI043222	Mm.42021	1431493
	Mus musculus mRNA for GM3 synthase,				
341	complete cds		AA274576.1		748275
	Mus musculus PAF acetylhydrolase				
342	mRNA, complete cds		AI324436.1		536464
242	Mus musculus pantophysin gene, complete		A A 071 FOF 1		625044
343	cds		AA271505.1		737944
211	Mus musculus rab6/rab5-associated protein (rab6) mRNA, partial cds		W77711.1		401958
344	Mus musculus skeletal muscle LIM		VV///11.1		401936
345	protein (FLH1) mRNA, complete cds		AA047966.1		477066
3.3	Mus musculus Stra13 mRNA, complete		12101750011		177000
346	cds		AA064241.1		480896
_	mWS3		In House-Clone		
		U35623,			
348	Myeloid cell leukemia sequence 1	AF063886	AA387843	Mm.1639	761106
349	Myostatin		AA052179.1		418993
	NAD(P)+ transhydrogenase (B-specific)				
	precursor	Z49204	AI323702	Mm.3842	580717
351	NADH dehydrogenase mwfe		W83104		404499
	NADH-ubiquinone oxidoreductase 13 kDa				
352	subunit	L38438	AA397301		599804

	NADH-ubiquinone oxidoreductase 13 kDa subunit	U59509	AA035972		468848
	NADP transhydrogenase	Z49204	AI323702	Mm.3842	580717
	Nitric Oxide Synthase-2		AA512708	Mm2893	922250
	Nitric Oxide Synthase-3		AA177240	Mm12837	620940
	Nuclear respiratory factor-1	AF098077	AI594316		1006311
	Nuclear respiratory factor-2	U20532		Mm.1025	635541
250	ornithine carbamoyltransferase precursor	X07092	AI266937	Mm.2611	1891345
339	ornitime carbamoyin ansierase precarse.	120.052	3220074.		
360	ornithineoxo-acid transaminase precursor	X64837	AI196410	Mm.13694	1887672
	р63а		In House-Clone		
	PACD		In House-Clone		•
	PEBP2		In House-Clone		
	Peripheral myelin protein, 22 kDa		AA416246.1		846064
304	empherat myemi protein, 22 -20 -				
365	peripheral-type benzodiazepine receptor 1	U12419	AA068577		524463
	Perox. Proliferator receptor (PPAR)				
366	Gamma	U01841	W34083	Mm.3020	317536
	Peroxisomal/Mitochondrial dienoyl-CoA				
367	isomerase		W29607		338088
		J03928,			
368	Phosphofructokinase 1	AF123533	AI480449	Mm.1166	862787
369	phospholipase A2, platelet, synovial fluid	X74266	AA871547	Mm.4675	1096251
370	phosphoprotein phosphatase	M81475	AI449151	Mm.1567	619279
371	PMP 35	L27842	AI573377	Mm.16453	534171
	probable aconitate hydratase,				
	mitochondrial (EST)	AI385870	AA275929	Mm.30065	775753
373	Procollagen Type 1		AA073604		536306
	Procollagen, type I, alpha 1		AA073604.1		536306
	Procollagen, type III, alpha 1		W89883.1		420322
376	Procollagen, type VI, alpha 1		W33786.1		352450
377	Protein Phosphatase inhibitor 2 (IPP2)	AA041826	AA041826	Mm.29617	475407
		U25114,			
	protoporphyrinogen oxidase	D45185		Mm.4730	482868
	pyruvate carboxylase	L09192	AI303529	Mm.1845	1888741
380	Pyruvate decarboxylase		AA308254		473778
381	Pyruvate dehydrogenase	M76727	AA423301		820409
382	pyruvate dehydrogenase (lipoamide)	M76728	AI323722	Mm.4223	513684
383	pyruvate dehydrogenase (lipoamide)	M76727	AA466268	Mm.34775	888842
	Pyruvate dehydrogenase E1 a subunit				
384	(human)	L13318	AA238899		888842
385	pyruvate kinase	D38379	AI035313	Mm.2635	1432851
	Pyruvate kinase	D63764	AI195164	Mm.8359	1886895
38	7 Pyruvate kinase	1	AA475121		873690
	Pyruvate kinase-like protein		W17814		334876
	RAB1		In House-Clone		ļ
	0 RAB1, member RAS oncogene family		AA175510.1		619501
	1 Rat NRBF1		AA259674		735186
	2 Rnase P-complex (RS Williams work)	U31228	AI614577		523232
	3 S100		In House-Clone		
_	4 sarc.		In House-Clone		
125	5 Sim to 6-Phosphofructo-2-kinase (human) M19938	AA397024		693346

Sim to acetyl CoA acetyl transferase (hum) D90228	1	į	1			
398 Sim to activel Coenzyme A synthetase AA37637 34949423 399 Sim to a-ketoglutarate (hum) D10523 W13320 329728 Min to arginyl-tRNA synthetase (Sac Cer) Min. 24457 423605 Min. 24363 576572 Min. 2363 576572 Min. 2363 576572 Min. 2363 576572 Min. 2363 Min. 2363 576572 Min. 2363 Min. 23660 Min. 24457 403660 Min. 24457 403660 Min. 24457 403660 Min. 2478 synthase sepsiton (Bov) AA108733 698109 Min. 2478	396	Sim to acetyl CoA acetyl transferase (hum)	D90228	AA272067		761668
399 Sim to a-ketoghutarate (hum)	397	Sim to acetyl Coenzyme A synthetase		AA109675		570550
Mm. 24457 423605	398	Sim to acetyl Coenzyme A synthetase		AA537637		949423
Min. 2363 576572	399	Sim to a-ketoglutarate (hum)	D10523	W13320		329728
Min. 2363 576572	400	Sim to Aldehyde dehydrogense (HUM)			Mm.24457	423605
402 Sim to ATP synthase epsilon (Bov)					Mm.22363	576572
Mathematical Color Mathema	402	Sim to ATP synthase epsilon (Bov)		AA108733		571214
Mathematical Color Mathema	403	Sim to ATP synthase F chain	PIR:A54211	W82194		403660
Sim to b-enolase (human) X56832 W11965 316967 Sim to branched-chain a-ketoacid dehydrogenase kinase TR:G924921 AA059497 480575			. X69908	AA239148		698109
Sim to branched-chain a-ketoacid dehydrogenase kinase Sim to carbon catabolite repressor prot.			X56832	W11965		316967
Sim to carbon catabolite repressor prot. Sim to carbon catabolite repressor prot. Sim to carbon catabolite repressor prot. Sim to carbon catabolite repressor prot.	1					
Sim to carbon catabolite repressor prot.	_		TR:G924921	AA059497		480575
May May	407	(Sac)		AA051133		438774
409 Sim to carnitine/acylearnitine carrier						
410 Sim to citrate synthase (Por) W14146 329884 411 Sim to cytochrome b5 (outer mito mem) X13923 AA203975 640762 412 Sim to cytochrome c oxidase VII X80899 AA050684 476180 Sim to electron trans flavoprotein a sub X71129 W18161 333641 414 (Hum) X71129 W18161 333641 415 Sim to electron trans flavoprotein b sub X71129 W18161 333641 416 Sim to enclase a subunit (Hum) M14328 AA204262 643854 417 Sim to Fructose 1-6-bisphosphate (Hum) L10320 AA276043 776124 418 Sim to glucose dehydrogenase (Bac. Sub) AA241896 680935 419 Sim to Glutamate/Malate trans (BOV) Mm.28466 582075 420 Sim to Glutamate/Malate trans (BOV) Mm.27395 317849 Sim to glycerol 3-phosphate W41175 351221 421 Sim to glycogen phosphorylase (Hum) J03544 W16286 334236 422 Sim to mito RNA pol (HUM) A1892781 Mm.34645 608625 Sim to Mito. 2-oxoglutarate/malate carrier (Human) AA245481 699237 Sim to NADH-ubiquin. oxidoreduct. 13 kd sub. AA245080 748891 Sim to NADH-ubiquin. oxidoreduct. 49 kd Sim to NADH-ubiquin. oxidoreduct. 9 kd sub. AA26232 871020 Sim to NADH-ubiquin. oxidoreduct. b14 sub. Sim to NADH-ubiquin. oxidoreduct. b15 Sim to NADH-ubiquin.						
Sim to cytochrome b5 (outer mito mem)	_					
Sim to cytochrome c oxidase VI B (Hum) X13923 AA139624 S81175						
Sim to cytochrome c oxidase VII						
Sim to electron trans flavoprotein a sub (Hum)	-	<u> </u>				
141	413		X80899	AA050684		476180
415 (Hum)	414	(Hum)	J04058	AA060723		481934
416 Sim to enolase a subunit (Hum) M14328 AA204262 643854 417 Sim to Fructose 1-6-bisphosphate (Hum) L10320 AA276043 776124 418 Sim to glucose dehydrogenase (Bac. Sub) AA241896 680935 419 Sim to Glutamate/Malate trans (BOV) Mm.28466 582075 420 Sim to Glutathione-S-transferase (RAT) Mm.27395 317849 Sim to glycerol 3-phosphate dehydrogenase 1 (rat) W41175 351221 421 Sim to glycogen phosphorylase (Hum) J03544 W16286 334236 423 Sim to hepatocyte gluc transporter AA002666 426758 424 Sim to mito RNA pol (HUM) A1892781 Mm.34645 608625 Sim to Mito. 2-oxoglutarate/malate carrier (Human) X66114 W54000 367801 426 Sim to mito. Elongation factor TS (Bov) AA245481 699237 Sim to NADH-ubiquin. oxidoreduct. 13 kd sub. AA288040 748891 Sim to NADH-ubiquin. oxidoreduct. 49 kd sub. AA321758 903911 430 Sim to NADH-ubiquin. oxidoreduct. ashl. W83085 404593 Sim to NADH-ubiquin. oxidoreduct. b14 sub. AA462323 871020 Sim to NADH-ubiquin. oxidoreduct. B14 Sim to NADH-ubiquin. oxidoreduct. B15 Sim to NADH-ubiquin. oxidoreduct. S15 Sim to NADH-ubiquin. oxidoreduct. S15 Sim to NADH-ubiquin. oxidoreduct. S16 Sim to NADH-ubiquin. oxidoreduct. S17 S172585 S18 to NADH-ubiquin. oxidoreduct. S18 S18 to NADH-ubiquin. oxidoreduct. S19 S18 to NADH-ubi			7771100	*****		
417 Sim to Fructose 1-6-bisphosphate (Hum) L10320 AA276043 776124 418 Sim to glucose dehydrogenase (Bac. Sub) AA241896 680935 419 Sim to Glutamate/Malate trans (BOV) Mm.28466 582075 420 Sim to Glutathione-S-transferase (RAT) Mm.27395 317849 Sim to glycerol 3-phosphate W41175 351221 421 dehydrogenase 1 (rat) W41175 351221 422 Sim to glycogen phosphorylase (Hum) J03544 W16286 334236 423 Sim to hepatocyte gluc transporter AA002666 426758 424 Sim to mito RNA pol (HUM) A1892781 Mm.34645 608625 Sim to Mito. 2-oxoglutarate/malate carrier (Human) X66114 W54000 367801 425 Sim to mito. Elongation factor TS (Bov) AA245481 699237 Sim to NADH-ubiquin. oxidoreduct. 13 kd sub. AA288040 748891 Sim to NADH-ubiquin. oxidoreduct. 49 kd sub. AA321758 903911 430 Sim to NADH-ubiquin. oxidoreduct. b14 sub. AA462323 871020	_	<u> </u>				
418 Sim to glucose dehydrogenase (Bac. Sub) AA241896 680935 419 Sim to Glutamate/Malate trans (BOV) Mm.28466 582075 420 Sim to Glutathione-S-transferase (RAT) Mm.27395 317849 Sim to glycerol 3-phosphate W41175 351221 421 dehydrogenase 1 (rat) W41175 351221 422 Sim to glycogen phosphorylase (Hum) J03544 W16286 334236 423 Sim to hepatocyte gluc transporter AA002666 426758 424 Sim to mito RNA pol (HUM) AI892781 Mm.34645 608625 Sim to Mito. 2-oxoglutarate/malate carrier X66114 W54000 367801 425 Sim to mito. Elongation factor TS (Bov) AA245481 699237 Sim to NADH-ubiquin. oxidoreduct. 13 kd AA288040 748891 Sim to NADH-ubiquin. oxidoreduct. 49 kd AA109715 572585 Sim to NADH-ubiquin. oxidoreduct. 9 kd AA521758 903911 430 Sim to NADH-ubiquin. oxidoreduct. ashl. W83085 404593 Sim to NADH-ubiquin. oxidoreduct. b14 AA462323 871020 Sim to NADH-ubiquin. oxidoreduct. W54068 367925						
419 Sim to Glutamate/Malate trans (BOV) Mm.28466 582075 420 Sim to Glutathione-S-transferase (RAT) Mm.27395 317849 Sim to glycerol 3-phosphate dehydrogenase 1 (rat) W41175 351221 422 Sim to glycogen phosphorylase (Hum) J03544 W16286 334236 423 Sim to hepatocyte gluc transporter AA002666 426758 424 Sim to mito RNA pol (HUM) AI892781 Mm.34645 608625 Sim to Mito. 2-oxoglutarate/malate carrier (Human) X66114 W54000 367801 426 Sim to mito. Elongation factor TS (Bov) AA245481 699237 Sim to NADH-ubiquin. oxidoreduct. 13 kd sub. AA288040 748891 Sim to NADH-ubiquin. oxidoreduct. 49 kd sub. AA109715 572585 Sim to NADH-ubiquin. oxidoreduct. 9 kd sub. AA521758 903911 430 Sim to NADH-ubiquin. oxidoreduct. b14 sub. AA462323 871020 Sim to NADH-ubiquin. oxidoreduct. b14 sub. AA462323 871020 Sim to NADH-ubiquin. oxidoreduct. b15 Sim to NADH-ubiquin. oxidoreduct. b15	417	Sim to Fructose 1-6-bisphosphate (Hum)	L10320	AA276043		776124
419 Sim to Glutamate/Malate trans (BOV) Mm.28466 582075 420 Sim to Glutathione-S-transferase (RAT) Mm.27395 317849 Sim to glycerol 3-phosphate dehydrogenase 1 (rat) W41175 351221 422 Sim to glycogen phosphorylase (Hum) J03544 W16286 334236 423 Sim to hepatocyte gluc transporter AA002666 426758 424 Sim to mito RNA pol (HUM) AI892781 Mm.34645 608625 Sim to Mito. 2-oxoglutarate/malate carrier (Human) X66114 W54000 367801 426 Sim to mito. Elongation factor TS (Bov) AA245481 699237 Sim to NADH-ubiquin. oxidoreduct. 13 kd sub. AA288040 748891 Sim to NADH-ubiquin. oxidoreduct. 49 kd sub. AA109715 572585 Sim to NADH-ubiquin. oxidoreduct. 9 kd sub. AA521758 903911 430 Sim to NADH-ubiquin. oxidoreduct. b14 sub. AA462323 871020 Sim to NADH-ubiquin. oxidoreduct. b14 sub. AA462323 871020 Sim to NADH-ubiquin. oxidoreduct. b15 Sim to NADH-ubiquin. oxidoreduct. b15	418	Sim to glucose dehydrogenase (Bac. Sub)		AA241896		680935
Sim to Glutathione-S-transferase (RAT) Sim to glycerol 3-phosphate					Mm 28466	
Sim to glycerol 3-phosphate dehydrogenase 1 (rat) W41175 351221						
421 dehydrogenase 1 (rat) W41175 351221 422 Sim to glycogen phosphorylase (Hum) J03544 W16286 334236 423 Sim to hepatocyte gluc transporter AA002666 426758 424 Sim to mito RNA pol (HUM) AI892781 Mm.34645 608625 Sim to Mito. 2-oxoglutarate/malate carrier (Human) X66114 W54000 367801 426 Sim to mito. Elongation factor TS (Bov) AA245481 699237 Sim to NADH-ubiquin. oxidoreduct. 13 kd AA288040 748891 Sim to NADH-ubiquin. oxidoreduct. 49 kd AA109715 572585 Sim to NADH-ubiquin. oxidoreduct. 9 kd AA521758 903911 430 Sim to NADH-ubiquin. oxidoreduct. b14 W83085 404593 Sim to NADH-ubiquin. oxidoreduct. b14 AA462323 871020 Sim to NADH-ubiquin. oxidoreduct. B14. Sub. W54068 367925 Sim to NADH-ubiquin. oxidoreduct. b15 W54068 367925						527015
422 Sim to glycogen phosphorylase (Hum) J03544 W16286 334236 423 Sim to hepatocyte gluc transporter AA002666 426758 424 Sim to mito RNA pol (HUM) AI892781 Mm.34645 608625 Sim to Mito. 2-oxoglutarate/malate carrier (Human) X66114 W54000 367801 425 Sim to mito. Elongation factor TS (Bov) AA245481 699237 Sim to NADH-ubiquin. oxidoreduct. 13 kd AA288040 748891 Sim to NADH-ubiquin. oxidoreduct. 49 kd AA109715 572585 Sim to NADH-ubiquin. oxidoreduct. 9 kd AA521758 903911 430 Sim to NADH-ubiquin. oxidoreduct. b14 AA462323 871020 Sim to NADH-ubiquin. oxidoreduct. b14 AA462323 871020 Sim to NADH-ubiquin. oxidoreduct. b15 W54068 367925	421			W41175		351221
423 Sim to hepatocyte gluc transporter AA002666 426758 424 Sim to mito RNA pol (HUM) AI892781 Mm.34645 608625 Sim to Mito. 2-oxoglutarate/malate carrier (Human) X66114 W54000 367801 425 Sim to mito. Elongation factor TS (Bov) AA245481 699237 Sim to NADH-ubiquin. oxidoreduct. 13 kd sub. AA288040 748891 427 Sim to NADH-ubiquin. oxidoreduct. 49 kd sub. AA109715 572585 Sim to NADH-ubiquin. oxidoreduct. 9 kd sub. AA521758 903911 430 Sim to NADH-ubiquin. oxidoreduct. ashl. W83085 404593 Sim to NADH-ubiquin. oxidoreduct. b14 sub. AA462323 871020 Sim to NADH-ubiquin. oxidoreduct. B14. Sub. W54068 367925 Sim to NADH-ubiquin. oxidoreduct. b15 W54068 367925			J03544			· · · · · · · · · · · · · · · · · · ·
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426 Sim to mito. Elongation factor TS (Bov) AA245481 699237 Sim to NADH-ubiquin. oxidoreduct. 13 kd AA288040 748891 Sim to NADH-ubiquin. oxidoreduct. 49 kd AA109715 572585 Sim to NADH-ubiquin. oxidoreduct. 9 kd AA521758 903911 430 Sim to NADH-ubiquin. oxidoreduct. ashl. W83085 404593 Sim to NADH-ubiquin. oxidoreduct. b14 AA462323 871020 Sim to NADH-ubiquin. oxidoreduct. W54068 367925 Sim to NADH-ubiquin. oxidoreduct. b15 W54068 367925					-	
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Sim to NADH-ubiquin. oxidoreduct. 49 kd AA109715 572585	127			A A 2880A0		749901
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430 Sim to NADH-ubiquin. oxidoreduct. ashl. W83085 404593 Sim to NADH-ubiquin. oxidoreduct. b14 AA462323 871020 Sim to NADH-ubiquin. oxidoreduct. W54068 367925 Sim to NADH-ubiquin. oxidoreduct. b15 W54068 367925	120	<u>-</u>		A A 521750		002011
Sim to NADH-ubiquin. oxidoreduct. b14 AA462323 871020						
Sim to NADH-ubiquin. oxidoreduct. 432 B14.5 sub. W54068 367925 Sim to NADH-ubiquin. oxidoreduct. b15		Sim to NADH-ubiquin. oxidoreduct. b14				
432 B14.5 sub. W54068 367925 Sim to NADH-ubiquin. oxidoreduct. b15	431			AA462323		871020
	432			W54068		367925
	433			AA434897		818906

1	Sim to NADH-ubiquin. oxidoreduct. b17	-		1]
434	sub.		W54448		367651
435			AA415725		846155
436			W83574		406509
437	Sim to NADH-ubiquin. oxidoreduct. Kfy1		W97248		423071
438	Sim to NADH-ubiquin. oxidoreduct. mn11 sub.		AA267638		723360
439	Sim to NADH-ubiquin. oxidoreduct. sgdh		AA222463		671212
440	Sim to NADH-ubiquin. oxidoreduct.15 kd		AA014507		439668
	Sim to putative mito. Ribosomal prot. S14		W89487		419614
442	Sim to quinone oxidoreductase		AA241313		653324
443	Sim to quinone oxidoreductase		AA259674		735186
	Sim to serine hydroxymethyltransferase, mito. (Rab)		AA208877		676311
445	Sim to sterol 26 hydroxylase			Mm.26793	578465
446	Sim to succinate dehydrogenase	M32246	AA108475		572339
	Sim to TIM17		W11644		318134
	Sim to Translocase of inner mito, mem. (Tim 44) (Sac)	Q01852	AA498767		888708
	Sim to ubiquinol-cty c reductase		AA087137		493604
	Sim to ubiquinol-cty c reductase, core prot.		AA108590		572127
451	Sim to UDP-glucose dehydrogenase		AA067191		523796
	Sim to UTP-Gluc1-phosphate uridyltrans (Human)	TR:G881394	AA473123		805218
	Sim to yeast ORF(Sim to rat synaptic glycoprotein) / SURF 4		AA220458		695279
	SKD 3		In House-Clone		<u> </u>
	Skel Mus LIM protein (FHL1) / RPB- assoc. mol. RAM14-1		AA047966		477066
	SMCK		In House-Clone		
	SMOO		In House-Clone		<u> </u>
	Solute carrier family 2	X15684	AA275871		776543
	Sperm capsule seleno protein	M88463	AA497767		917403
	sperm mitochondrial capsule selenoprotein	S49657, M29603	AI482284	Mm.554	917537
	steroidogenic acute reg protein precursor, mito	L36062	AA389406	Mm.3436	569013
	2 sterol carrier protein 2 precursor	M62361		Mm.1779	580813
	3 Succinate dehydrogenase-Cyt B560	TR:G1019861	AA137762		580040
	4 superoxide dismutase (Mn) precursor	Z18857, L35525	AA415267	Mm.2597	791140
	5 Superoxide dismutase-1 (Cu/Zn Sod)		AA039044	Mm5274	474545
	Superoxide dismutase-3 (extracellular 6 Sod)		AI314465	Mm2407	1907770
	Suppressor of Ty 6 (S. cerevisiae) 7 homolog		AI323032.1		477535
	8 Surfeit gene 4		AA220458.1		695279

469	Surfeit locus protein 1	M14689	AA274488	Mm.6874	748268
	tetrahydrofolylpolyglutamate synthase		•		
470	precursor	U32197	AA030778	Mm.3830	463571
471	Thioredoxin		AA242573		681159
472	Thioredoxin			Mm.3533	579774
473	Thioredoxin Reductase		AI529082	Mm 25543	1887250
474	thiosulfate sulfurtransferase	U35741	AI196763	Mm.15312	1887427
475	Thiosulfate sulfurtransferase (mito) Tst1	U35741	AI195057	Mm.15312	188763
476	TIMP2		In House-Clone		
477	Tissue inhibitor of metalloproteinase 2		AA444490.1		831964
478	Tissue inhibitor of metalloproteinase 2		AA518165.1		902923
479	TOAD		In House-Clone		
480	transforming protein bcl-2-alpha	L31532	AA867214	Mm.5155	1265430
	transforming protein bcl-2-beta	M16506	AA867214	•	1265430
482	transforming protein bcl-w (3' seq)	AF030769.1	AA563148		975210
483	transforming protein bcl-w (5'-seq)	AF030769.1	AA667328		1139352
484	Transketolase (Tkt)	U05809	AI132421	Mm.9307	1481358
485	Type II Peroxiredoxin 1		W83228	Mm42948	405943
486	Uncoupling Prot homology (UCPH)		AA260521		748122
487	Uncoupling Protein 2			Mm.12556	423616
488	Uncoupling protein 3		AA062091		482847
489	Unknown EST				608265
490	uracil-DNA glycosylase	U55040	W48179		355462
491	voltage-dependent anion channel 1	U230840	AA244874		680076
492	voltage-dependent anion channel 2	U30838	AI507203	Mm.569	931442
493	voltage-dependent anion channel 3	U30839	AA616007	Mm.38513	1066900

Example 3

Identification of Mutations Causing Disease

The mitochondrial respiratory complex I is assembled from seven mtDNA genes and thirty-six nDNA genes. Patients with complex I defects have phenotypes ranging from midlife-onset optic atrophy to lethal childhood Leigh's disease. Mitochondrial biology expression profiles were determined for patients with a variety of complex I defects. Samples are collected from a variety of patients with complex I defects. Each sample is reverse transcribed, labeled, and hybridized, together with standard target, to a human array comprising probes selected from Example 1. The hybridization measurements are analyzed, leading to the identification of several novel mtDNA mutations and dominant and recessive nDNA mutations.

Example 4

Profile for Complex IV Leigh's Syndrome

The mitochondrial biology expression profile was determined for a complex IV Leigh's syndrome (LS) patient. LS is a subacute neurodegenerative condition characterized

by necrotic lesions in the brain stem, basal ganglia, thalamus and spinal cord. Death is usually within 2 years of onset of symptoms that may include motor and/or intellectual retardation, abnormal breathing rhythm, nystagmus, opthalmoparesis, optic atrophy, ataxia, and dystonia. The Leigh's syndrome patient had a typical complex IV cytochrome c oxidase deficiency associated with surfeit 1 (SURF-1) gene mutations. This patient was from a consanguineous marriage and was homozygous for a nonsense mutation in the SURF-1 gene. Expression profiling of muscle and cultured cell samples from this patient using a human array of Example 1 was performed, in comparison to a control reference standard. NDUFS8 expression was not significantly altered. However, many nuclear and mitochondrially encoded complex I genes were down-regulated, including mtDNA transcripts ND4, NDL4, and ND6. Nuclear genes SURF-1, SOD2, 70kD heat shock protein, voltage dependent anion channel (VDAC4), adenine nucleotide translocase 2 (ANT2), and glutathione peroxidase 3 were down-regulated.

Example 5

Profile for Complex I Leigh's Syndrome

Mitochondrial biology expression profiles were determined for twelve complex I Leigh's syndrome patients (Procaccio, VF (2001) EuroMit5 Abstract). Sequencing of all 43 genes known to be part of complex I, of each patient, identified one patient as a compound heterozygote for two missense mutations in the 23 kD NADH dehydrogenase (NDUFS8) gene of complex I. This patient had a respiratory complex I defect apparent in skeletal muscle and cultured lymphoblastoid cells. Samples were collected from cultured lymphoblastoid cells from this patient and control reference lymphoblastoid cells. Samples were reverse transcribed and differentially labeled and hybridized to a human array comprising probes selected from Example 1. The expression profile was determined using a hierarchical clustering method. Mitochondrial biology expression profiles from the other patients were similarly determined using appropriate samples and controls. Expression profiles of all patients were characteristic of complex I deficiencies, including downregulation of all mtDNA and some nDNA complex I genes and up-regulation of the adenine nucleotide translocator genes (ANT1 and ANT2).

Example 6

Diagnosis of Complex IV Leigh's Syndrome

The mitochondrial biology expression profile for Leigh's syndrome SURF-1 nonsense mutations, as determined in Example 4, is used to diagnose patients. Samples are collected from patients and mitochondrial biology expression microarray-tested using a human array containing probes for at least SURF-1, ND4, NDL4, ND6, SOD2, 70kD heat shock protein, VDAC4, ANT2, and glutathione peroxidase 3.

Example 7

A Mouse MitoChip

A mouse Mitochip was printed with probes for 452 genes. Some of these genes were represented by two or more probes, providing internal controls for the reproducibility of gene expression quantitation. An additional 37 control spots were included on the array. Of these, 25 were probes for housekeeping genes to allow normalization between samples. The remaining 12 spots were various controls for hybridization and positioning. Table 2 lists the functional categories and number for all of the housekeeping genes on this array. The cDNA clones that represent each gene were either from the I.M.A.G.E. consortium or cloned by The Center for Molecular Medicine and published in (Murdock et al., 1999). A complete annotation of each gene was compiled and GenBank accession numbers and Unigene cluster numbers were determined. Table 5 provides a list of the probes on this array.

Example 8

Profile of Sod2 Heterozygote Mutant Mice at Various Ages

Oxidative stress has been implicated in aging and degenerative disease. Mitochondria are thought to be the main source of reactive oxygen species such as superoxide anion. Mitochondrial superoxide anion is normally detoxified by manganese superoxide dismustase (MnSOD, the Sod2 gene). However, when, free radical metabolism is perturbed, oxidative damage to protein, DNA, and lipids may occur. To demonstrate the effects of increased superoxide anion toxicity on mitochondrial physiology with age, the mitochondrial biology expression profiles of mice with a 50% reduction in MnSOD (Sod2 +/-) were determined at various ages. Samples were collected from young (5 months), middle-aged (10-14 months), and old (20-25 months) wild-type and Sod2 +/- mice. Samples were reverse transcribed and differentially labeled from the corresponding controls. The labeled mutant sample and the corresponding labeled control were hybridized with the mouse array of Example 2. Relative

to the control mice, the old Sod2 +/- mice showed induction of antioxidant and apoptosis genes including glutathione peroxidase 3, apoptosis inhibitory factor 3, caspase 1, and the peripheral benzodiazepine receptor.

Example 9

Profile of Sod2 Homozygote Mutant Mice

Manganese superoxide dismutase (MnSOD, the Sod2 gene) is a gene expression product involved in mitochondrial biology. Sod2 -/- animals die soon after birth due to the superoxide inactivation of mitochondrial iron-sulfur center enzymes resulting in dilated cardiomyopathy. The mitochondrial biology expression profile of Sod2 -/- mice is determined using the mouse MitoChip of Example 2. RNA samples are collected from Sod2 -/- mice and Sod2 +/+ mice. The Sod2 -/- sample is reverse transcribed and labeled with Cy3 phosphoramidite. The Sod2 +/+ sample is reverse transcribed and labeled with Cy5 phosphoramidite. The labeled samples are incubated with a mouse array under conditions of high stringency hybridization. The hybridization of both samples is measured with a microarray reader. The hybridization measurements are recorded.

Example 10

Profile of GPxI Mutant Mice

Glutathione peroxidase 1 (GPx1) is an expressed sequence involved in mitochondrial biology. GPx1 -/- animals show mild growth inhibition and reduced OXPHOS efficiency. The mitochondrial biology expression profile of GPx1 -/- mice is determined using a mouse array of Example 2. RNA samples are collected from GPx1 -/- mice and GPx1 +/+ mice. The GPx1 -/- sample is reverse transcribed and labeled with Cy3 phosphoramidite. The GPx1 +/+ sample is reverse transcribed and labeled with Cy5 phosphoramidite. The labeled samples are incubated with a mouse array under conditions of high stringency hybridization. The hybridization of both samples is measured with a microarray reader. The hybridization measurements are recorded.

Example 11

Profile of Sod2 Heterozygote GPx1 Homozygote Doubly Mutant Mice

The mitochondrial biology expression profile of Sod2 -/+ plus GPx1 -/- mice is determined using a mouse array of Example 2. RNA samples are collected from Sod2 -/+ plus GPx1 -/- mice and Sod2 +/+ plus GPx1 +/+ mice. The Sod2 -/+ plus GPx1 -/- sample is

reverse transcribed and labeled with Cy3 phosphoramidite. The Sod2 +/+ plus GPx1 +/+ sample is reverse transcribed and labeled with Cy5 phosphoramidite. The labeled samples are incubated with a mouse array under conditions of high stringency hybridization. The hybridization of both samples is measured with a microarray reader. The hybridization measurements are recorded.

Example 12

Profile of Mutant Mice Overexpressing Sod2 and/or GPx1

The mitochondrial biology expression profiles are determined using a mouse array, for mice overexpressing MnSOD and for mice overexpressing MnSOD plus GPx1.

Example 13

Profile of ρ⁰ Mutant Cell Line

A mouse array of Example 2 was used to determine the mitochondrial biology expression profile of the mouse mutant cell line ρ^0 , the most extreme case of mitochondrial dysfunction. The LMEB4 (ρ^0) cell line was profiled against its parental LM(TK) - cell line. The mouse mutant cell line ρ^0 lacks mitochondrial DNA. To maintain the LMEB4 cell line in culture, it must be grown in media supplemented with glucose, pyruvate, and uridine (GUP media). A scatter plot of the gene expression ratios is shown in FIG. 3. Samples from the ρ^0 cell line and from the LM(TK) cell line were reverse transcribed and differentially labeled using a standard two-color fluorescent system, and hybridized to a mouse array of Example 2. Mouse array analysis confirmed that all mtDNA-encoded transcripts were absent from the LMEB4 cells, and that there was a reduction in nDNA OXPHOS gene expression, aconitase, and nuclear receptor binding factor 1 (NRBF1). There was an increase in expression of key glycolytic genes, mitochondrial ribosomal proteins, the LON protease, heat shock protein 84 (HSP 84), Bcl-X binding protein, and antioxidant protein 1. Invariably, the nuclear-encoded OXPHOS complex subunits were also down-regulated between 3 and 38-fold with a mean of 4.5 (the mean was calculated excluding the single outlying complex I subunit NADHdehydrogenase mwfe which was down-regulated 38-fold). Mitochondrial transport proteins such as the Glutamate-malate transporter were down-regulated as was the mitochondrial protein import subunit gene Tim17 and several amino acid metabolism genes. By contrast, glycolytic genes such as pyruvate kinase, glucose phosphate isomerase and glucose-6phosphate dehydrogenase were up-regulated 2 to 3-fold. Phosphofructokinase was up 1.6fold. Anti-apoptotic genes such as apoptosis inhibitor 2 and 3 were up-regulated as was the

pro-apoptotic Bcl-Xs binding protein BNIP3 and Caspase 2. The other Bcl protein family members that are on the array were not changed significantly. The multi-function mitochondrial LON protease was up-regulated 2.1-fold.

Example 14

Profile of CAPR Mutant Cell Line

A mouse array of Example 2 was used to determine the mitochondrial biology expression profile of the mouse mutant cell line harboring a mutation for chloramphenicol resistance (CAPR), and the CAPR 501-1 cell line having a mtDNA mutation in the 16S rRNA The CAP^R mutation in chimeric mice causes cataracts, reduced photoreceptor response, vacuolization of the retinal pigment epithelium, and hamartomatous outgrowths of the optic nerve head. Mice inheriting the CAPR mutation showed a marked increase in embryonic lethality, and those that were born died within two weeks with growth retardation, dilated cardiomyopathy, and mitochondrial abnormalities. CAPR 501-1 was compared to the CAPS LM(TK) - cell line. These two cell lines are both derived from mouse L929 cells. Samples from the CAPR cell line and from wild-type cells were reverse transcribed and differentially labeled with a standard two-color fluorescent system, and hybridized to a mouse array of Example 2. The CAPR cell line had up-regulation of all thirteen mtDNA transcripts, but down-regulation of multiple nDNA OXPHOS genes. The CAPR 501-1 cell line versus the LM(TK)- gene expression scatter plot showed that all mtDNA transcripts were up-regulated 3.1 to 3.5-fold while the nuclear encoded OXPHOS subunits were downregulated 2.1 to 5.3-fold. Procollagen type III and VI were also up-regulated 3.5 to 4-fold.

Example 15

Profile of Treatment to Cell Line

Mouse arrays of this invention were used to demonstrate how treatment changes, such as changing cell culture conditions, affect gene expression. The control cell line LM(TK) - grown in standard medium was profiled against a culture of LM(TK) - cells grown in media supplemented with glucose, pyruvate, and uridine (LM(TK) - (GUP)). Samples from the treated fibroblast cell line and from untreated fibroblast cells were reverse transcribed and differentially labeled with a standard two-color fluorescent system, and hybridized to a mouse array of Example 2. Treatment resulted in a down-regulation of the LON protease and HSP 84. The scatter plot of this experiment showed that other than the same core group of genes that were up-regulated in the NZB cell line mentioned in Example 17, few genes were

significantly different in their expression. The hybridization spots of three genes that showed the highest differences were HSP70, the LON protease, and E.T.F. The 70 kDa heat shock protein (HSP70) was down-regulated 3.4-fold. HSP70 has been shown to be a chaperone protein involved in mitochondrial protein import that forms an ATP-dependent motor with the inner mitochondrial membrane translocase and the polypeptide in transit (Voos, W. et al., "Mechanisms of protein translocation into mitochondria," [1999] Biochimica et Biophysica Acta 1422:235-54). The entire HSP70 control spot was of medium intensity, while the experimental spot was only medium intensity in the center. The LON protease was downregulated 9.7 fold in LM(TK) - cells grown in GUP. The control LON protease spot was of medium high intensity over the entire spot and of low intensity in the experimental spot. The electron transfer flavoprotein (ETF), which shuttles electrons gathered during fatty acid metabolism to the electron transport chain, was down-regulated 3.8 fold. The E.T.F control spot was high intensity and the experimental spot very low intensity. Some of the nuclear encoded OXPHOS subunits as well as several proteins involved in amino acid metabolism were down-regulated 1.5 to 2-fold with mean ratio of 1.65. Since most of these genes fell below the +/- 1.7 ratio cutoff, further analysis was needed to determine if the expression pattern was significant. There were no differences in mtDNA transcript levels and no consistent pattern of up-regulation of glycolytic genes.

Example 16

Profile of Sod2 Mutant Mice After Treatment and Before Symptoms

Treatment of Sod2 mutant mice with MnTBAP prevents cardiac and liver pathology, however after 12 days the MnTBAP-treated mutant animals develop a prominent movement disorder which leads to debilitation by three weeks, in association with spongiform changes and gliosis in the cortex and specific brain stem nuclei associated with motor function. It is thought that the severe neuropathology results from poor exchange of MnTBAP across the blood brain barrier. The mitochondrial biology expression profile of MnTBAP-untreated, Sod2 mutant mice and MnTBAP-treated, Sod2 mutant mice was determined using the mouse array of Example 2. Samples were collected from 8 day old Sod2 mice without MnTBAP treatment, 8 day old Sod2 mice with MnTBAP treatment, and 12 day old Sod2 mice with MnTBAP treatment. Samples were also collected from age-matched controls. About 20 genes were found to be differentially expressed in all three groups of Sod2 knockout mice compared to the corresponding age-matched controls. The about 20 genes included bioenergetic genes such as the mitochondrial creatine phosphokinase, antioxidant enzymes

like the glutathione peroxidase 3, and apoptotic factors including caspase 1 and apoptosis inhibitor factor 3. The excitatory amino acid transporter 3, frataxin, and one EST of unknown function were also induced. Mitochondrial biology expression profiling demonstrated changes in expression before neuropathic changes were manifested.

Example 17

Organ-Specific Profiles of Mutant Mice

The NZB mouse line mtDNA and the "common haplotype" mtDNAs (129/Sv, C57Bl/6J, C3H, BALB/c, and others which are thought to have arisen as the progeny of a single female (Ferris et al.,1982) differ by 108 nucleotides, and these polymorphic differences have been used to monitor the segregation of heteroplasmic populations of mtDNAs in mice created by embryo fusion techniques (Jenuth, J. P. et al., "Random genetic drift in the female germline explains the rapid segregation of mammalian mitochondrial DNA," [1996] Nat Genet 14:146-51; Jenuth, J. P. et al., "Tissue-specific selection for different mtDNA genotypes in heteroplasmic mice," (1997) Nat Genet 16:93-5; Meirelles, F.V., and Smith, L.C., "Mitochondrial genotype segregation in a mouse heteroplasmic lineage produced by embryonic karyoplast transplantation," (1997) Genetics 145:445-51; Meirelles, F.V. and Smith, L.C., "Mitochondrial genotype segregation during preimplantation development in mouse heteroplasmic embryos," [1998] Genetics 148:877-83). Tissues from the NZB and CAPR mice were profiled on a mouse array. Messenger RNA was isolated from the brain, liver, spleen, kidney, heart, and skeletal muscle of a male mouse heteroplasmic for the NZB mtDNA and a male mouse that was 80% chimeric for ES cell-derived CAPR cells as defined by coat color. Due to the severity of the CAPR mutation it was not possible to analyze the mitochondrial gene expression changes in mice that were homoplasmic for the CAPR mtDNA. Control mRNA for each of the tissue samples was isolated from sex, age, and nuclear background-matched control mice. All of the tissue samples were genotyped to determine the levels of heteroplasmy for the NZB and CAPR mtDNA in each of the tissues. Equal levels of the NZB and "common" mtDNA were found in the six tissues analyzed from the NZB mtDNA-positive mice. The six tissues from the CAPR chimera had varying levels of CAPR mtDNA with the kidney and spleen having the highest amounts, 65% and 50% CAPR mtDNA, respectively. The heart contained approximately 20% CAPR mtDNA, while brain, liver, and muscle all contained between 5% and 10% CAPR mtDNA. Analysis of the NZB-mtDNA tissue samples did not reveal any differentially expressed genes in the heart, liver, brain, and kidney. A scatter plot from the NZB heart is shown in FIG 4. The scatter

plots from the liver, brain, and kidney are virtually identical in that nearly every gene has an expression ratio of 1. Analysis of the NZB-mtDNA spleen and muscle showed several genes that were differentially expressed in the two tissues. The NZB-mtDNA muscle showed a 1.5 to 2.1-fold reduction in all mtDNA transcripts, pyruvate dehydrogenase was down 2.2-fold, and there was a general trend for nuclear-encoded OXPHOS subunits to be down-regulated 1.4 to 1.8-fold. The vesicular transport protein, pantophysin, was down-regulated 4-fold and the glycogenolysis rate-limiting enzyme, glycogen phosphorylase, was down 3-fold. There were not any genes that were significantly up-regulated in the muscle. A similar pattern of mtDNA-encoded gene expression was observed in the NZB-mtDNA spleen with all transcripts down 1.8 to 2-fold. However, there were no differences in nuclear OXPHOS subunit expression levels like that observed in the NZB-mtDNA skeletal muscle. In contrast to the NZB-mtDNA muscle, several genes were up-regulated in the spleen. opposition to the results in the NZB cell line, both probes of the heme biosynthesis gene coproporphyrinogen oxidase III derected up-regulation 3-fold in the spleen. The integral membrane protein SURF 4 was up 2-fold and the amino acid metabolism gene 2-amino-3ketobutyrate CoA ligase was up 4.8-fold. Glycogen phosphorylase, down 3-fold in the muscle, was up 4.8 fold in the spleen. The muscle and spleen results suggest that the polymorphisms between the NZB and "common" mtDNA may have a functional consequence in some tissues but not others. Analysis of the CAPR tissue samples did not show any genes to be differentially expressed in the kidney, heart, muscle, liver, or spleen. The kidney, having the highest percentage of mutant mtDNAs, had expression ratios around 1 for nearly every gene. The two outliers on the kidney scatter plot that appear to be downregulated can be explained by hybridization artifacts causing a high background in the control sample. The CAPR brain sample was the only tissue that had any differentially expressed Skd 3 was up-regulated 2.2-fold, glutathione peroxidase was up 2.4-fold and apoptosis-inhibitor 3 was up 2.4-fold. Although no genes were down-regulated in the brain more than 1.8-fold, closer analysis of the brain samples did reveal a trend that was not observed in any of the other tissues. Several nuclear-encoded OXPHOS subunits were downregulated between 1.3 and 1.6-fold. These included five Complex I subunits, three Complex IV subunits and five Complex V subunits as well as VDAC 1 and 3. None of the Complex II and III subunits or mtDNA transcripts followed this trend. Principal component analysis of NZB and CAP^R mouse tissues, separately and together with the cell lines, was performed.

Example 18

Identification of Genes for Mitochondrial Arrays

Mice mutant in mitochondrial biology were used to identify genes involved in mitochondrial biology. Mice deficient in the heart/muscle isoform of the adenine nucleotide translocator (ANT1) exhibit many hallmarks of human oxidative phosphorylation (OXPHOS) disease, including dramatic proliferation of skeletal mitochondria. Samples were collected from the gastrocnemius muscle of ANT1 and wild-type mice, reverse transcribed and differentially labeled, and hybridized with a mouse microarray chip (Mouse Unigene 1, Incyte Genomics Inc., Palo Alto, California) containing over 8000 sequence-verified cDNAs. Analysis of the hybridization results identified more than 150 differentially expressed genes. Gene sequences that were not previously recognized as being involved in mitochondrial biology were used to generate probes that were placed on the mouse array of Example 2. Homologous human gene sequences were used to generate probes that were placed on the human array of Example 1.

Example 19

Profile of Age-Related Changes in Chimpanzee Using Human Mitochondrial Array

Age-related changes in the mitochondrial biology expression profile in chimpanzees are determined using a human array of Example 1. Samples from young adult chimpanzee muscle and samples from most-mortem tissues of older chimps are reverse-transcribed, differentially labeled, and hybridized with a human array of Example 1.

Example 20

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Profile of Putative Neutral Variant Mutant Mouse

The NZB cell line was profiled to examine the changes in mitochondrial gene expression resulting from a more neutral set of mtDNA polymorphisms. As mentioned previously, the NZB mtDNA contains 108 sequence differences compared to the "common" mouse mtDNA genotype found in LM(TK). While these differences were reported to be neutrally polymorphic (Jenuth et al., [1996] Nature Genetics 14:146-151; Meirelles and Smith [1997] Genetics 145:445-451), the only evidence to support that hypothesis is that transgenic mice containing a high percentage of NZB mitochondria have no overt phenotypes (Levy, S. E., "Genetic Alteration of the Mouse Mitochondrial Genome and Effects on Gene Expression," (2000) Ph.D. Thesis, Emory University; Jenuth et al. [1997] Nature Genetics 16:93-95; Meirelles and Smith [1998] Genetics 148:877-883). An NZB cybrid cell line was

W₀ 03/020220 PCT/US02/27886

profiled on a mouse mitochondrial array. The scatter plot of gene expression ratios between the NZB cell line and the parental LM(TK) - (without GUP supplementation) shows that both probes of the fatty acid metabolism gene Acyl-CoA dehydrogenase (medium-chain) detected up-regulation 3.6-fold. Procollagen III and VI were up-regulated 6.2 and 6.8-fold, respectively. Two independent probes of the coproporphyrinogen oxidase III gene that is involved in heme biosynthesis detected down regulation 2.6 and 2.3-fold. Also downregulated was the peripheral-type benzodiazepine receptor. This receptor has been implicated in a variety of mitochondrial functions including the regulation of mitochondrial protein import under conditions of oxidative stress, calcium homeostasis, and steroidogenesis (Culty, M. et al., "In vitro studies on the role of the peripheral-type benzodiazepine receptor in steroidogenesis," [1999] J. Steroid Biochemistry & Molecular Biology 69:123-30; Wright, G., and Reichenbecher, V. "The effects of superoxide and the peripheral benzodiazepine receptor ligands on the mitochondrial processing of manganese-dependent superoxide dismutase," [1999] Experimental Cell Research 246:443-50). The glycolytic genes glyceraldehyde-3-phosphate dehydrogenase and glucose-6-phosphate isomerase were upregulated 1.7 and 2.1-fold, respectively. Glycolytic genes were also up-regulated in the NZB cell line. This indicates that the sequence polymorphisms between the NZB and "common" mtDNAs are not entirely neutral and cause changes in mitochondrial function when combined with the LM(TK) - nucleus. Thus, the NZB mtDNA does not appear to be completely interchangeable with the "common" mtDNA genome. An interesting group of genes that were up-regulated in the NZB cell line were the pro-inflammatory genes Caspase 1 and platelet activating factor (PAF) acetylhydrolase, the mitochondrial RNA polymerase, and glutathione peroxidase 3.

Example 21

Hierarchical Clustering of Cell Lines

Principal component analysis (PCA) and hierarchical clustering were performed on the cell line data (Examples 13-15 and 20) to group genes based on similarities in their expression patterns over all the samples. PCA analysis was used to reduce the dimensionality of the data by calculating three principal axes that encompass as much of the variability in all of the samples as possible. Each of the samples was then plotted on those axes in three-dimensional space. The PCA results revealed that the NZB cell line clustered away from the other cell lines, consistent with it having fewer differentially expressed genes in common with the other samples. The LMEB4 $\rho 0$, 501-1 and LM(TK) - (GUP) cell lines all arrayed

along one common principle axis, probably due to the commonality of a down-regulation of nuclear OXPHOS genes. However, they were divergent in the other two axes. The LM(TK) - (GUP) and NZB did share one axis, possibly due to a partial reduction in OXPHOS genes and a concomitant induction of glycolytic gene expression. However, both showed few differences when compared to the CAP R 501-1 and LMEB4 p0 samples. A hierarchical clustering algorithm was used to group genes with similar expression profiles across all of the samples. Both genes as well as samples were clustered together using a Euclidean distance measurement and average linkage. The clustering results revealed seven groups of genes with similar expression patterns in the cell line samples. Certain classes of genes were found to change together. Similar to the PCA analysis, the expression changes seen in the LM(TK) -(GUP) and NZB samples clustered closest together with the CAP R 501-1 and LMEB4 ρ0 samples branching successively further away. The Group 1 genes are involved in fatty acid metabolism. Group 2 genes, mainly down-regulated in the LM(TK) - (GUP) sample, include malate dehydrogenase, lactate dehydrogenase, glucose phosphate isomerase, and several amino acid metabolism genes. Group 3 genes are diverse clusters of genes that change in expression coordinately across the 5 samples. It includes some nuclear-encoded OXPHOS subunits, a few antioxidant and transport proteins as well as pyruvate kinase and a GTPbinding protein. Group 4 is a small, diverse cluster of genes mainly up-regulated in the CAP R 501-1 cell line. This group includes several of the same genes found to be up-regulated in the Antl(-/-) mouse by differential display analysis (Murdock et al., 1999). Also in this group are two NADP-transhydrogenases, carbonate dehydratase and cytochrome b5 reductase. Group 5 is almost exclusively nuclear-encoded OXPHOS subunits. The voltage-dependent anion channel (VDAC) genes and several antioxidant proteins also cluster in this group. Group 6 is composed almost entirely of mtDNA-encoded transcripts. Group 7 is the heterogeneous group of genes that were up-regulated in each of the samples analyzed and included Caspase 1, PAF acetylhydrolase, the mitochondrial RNA polymerase, and glutathione peroxidase 3. Hierarchical clustering packages are available in the art, i.e. Expression Profiler (http://ep.ebi.ac.uk/EP/ from the European Bioinformatics Institute, Cambridge, UK). PCA is described in Bioinformatics 2001, volume 17, number 9, pages 763-774.

The following examples describe the preparation of a mitochondrial biology expression array, sample preparation, hybridization, scanning, and data normalization.

Example 22

PCR Amplification

PCR amplifications were performed with standard PCR techniques. Probes were made my amplifying clones using a universal primer set (Forward primer 5'-CTGCAAGGCG ATTAAGTTGGGTAAC-3' Reverse primer 5'-GTGAGCGGATAACAATTTCAC ACAGGAAACAGC-3') in a 100 μl PCR reaction containing PCR buffer (10 mM Tris, 1.5 mM MgCl2, 50 mM KCl, pH8.3), 0.2 mM dNTPs, 0.2 mM each primer, 1.25 U Taq (Sigma, St Louis, MO). 0.5-1 μl of bacterial culture was added to each PCR reaction and thermal cycling was done as follows: 4 minutes at 94 C followed by 35 cycles of 15 seconds at 94 C, 30 seconds at 66 C and 1 minute 30 seconds at 72 C. Following cycling, reactions were held at 72 C for 4 minutes to complete all extension reactions. All PCR products were confirmed by agarose gel electrophoresis through a 1.5% gel. After satisfactory amplification, products were quantitated by UV 260/280 ratio and desiccated in a Savant Speed-Vac (Holbrook, NY). Dried products were then resuspended in 3xSSC (450 mM NaCl, 40 mM sodium citrate) at a concentration of 400-600 ng/μl for arraying.

Example 23

Slide Preparation

Before arraying probes, the glass microscope slides for the arrays were coated with poly-Lysine to provide a substrate for DNA binding. Standard glass microscope slides (Gold Seal, Beckton-Dickson, Franklin Lakes, NJ) were cleaned in a solution of 2.5 M NaOH, 60% ethanol for two hours. After cleaning, slides were rinsed five times in fresh water. The slides were then soaked in a solution of 0.01% poly-L-lysine, .1x PBS for 1 hour followed by rinsing in fresh water. After rinsing, the slides were dried in a vacuum oven at 45°C for 15 minutes.

Example 24

Printing

Arrays were printed onto poly-L-lysine coated glass slides using the GMS 417 Arrayer (Affymetrix/Genetic Microsystems, Woburn, MA). The arrays were printed using a 4-pin print head with a spot size of 150 μ m (approximately 33 pL of volume per spot) and a center-to-center spot spacing of 375 μ m. A humidity level of 65-70% was maintained during the printing of the arrays by a custom humidifier system. After printing, the arrays were

allowed to dry for 1 hour at room temperature. The arrays were then processed by rehydrating over a warm solution of 1x SSC for 5 minutes followed by rapid drying on a 95°C heat block. Following drying, the DNA was crosslinked to the slide by exposing the arrays to 65 mJ of ultraviolet energy (Stratalinker, Stratagene, La Jolla, CA). To block non-specific interactions on the arrays during hybridization, the slides were then treated with a solution of 60 mM succinic anhydride and 40 mM sodium borate in 1-methyl-2-pyrrolidinone for 15 minutes at room temperature. The arrays were then denatured in 95°C water for 2 minutes and dehydrated by rapid immersion in 95% ethanol. The arrays were then dried by centrifugation at 20xg for 5 minutes.

Example 25

Sample Preparation

Total RNA preparations were performed using the TRIzol reagent (Life Technologies, Gaithersburg, MD) as per the manufacture's directions. For cell culture samples, a 90% confluent 225ml flask was lysed directly in the flask with 18 ml of TRIzol. At least three flasks were pooled for each cell line to reduce any variability caused by culture conditions. For each mouse tissue, RNA was isolated from approximately 500 mg of tissue that was mechanically homogenized in 6ml of TRIzol. Following the isolation of total RNA, poly-A+mRNA was isolated using Qiagen Oligotex (Valencia, CA) as per the manufacture's directions.

Example 26

Reverse Translation, Labeling, and Hybridization

To produce targets for hybridization to the MitoChip arrays, 2 μg of poly-A+ RNA was labeled with fluorescent nucleotides by reverse transcription. The poly-A+ RNA was mixed with 3 mg of anchored oligo-dT and incubated at 70°C for 10 minutes followed by 10 minutes on ice. The denatured and annealed RNA was then reverse transcribed in a 30 μl reaction mix containing reaction buffer (50 mM Tris-HCl, 75 mM KCl, 3 mM MgCl2 pH 8.3), 10 mM dithio-threatol, 500 μM dATP,dGTP,dTTP, 300 μM dCTP, 20 U SuperScript reverse transcriptase (Life Technologies, Gaithersburg, MD) and 100 μM of either Cy5-dCTP (control samples) or Cy3-dCTP (experimental samples). The reactions were incubated at 42°C for 2 hours. Following incubation, 15 μl of 0.1M NaOH was added to degrade the remaining template RNA and the sample incubated at 70°C for 10 minutes. The reaction was neutralized by the addition of 15 μl of 0.1 M HCl followed by 440 μl of TE buffer (10 mM

Tris, 1 mM EDTA, pH 7.4). The synthesized cDNA was purified by size-exclusion filtration using Microcon YM-3 centrifugal filter devices (Millipore, Bedford, MA). After purification, 10 μg of poly-A RNA (Sigma, St Louis, MO) and 10 μg of yeast transfer RNA (tRNA) (Life Technologies, Gaithersburg, MD) was added. The final sample volume was adjusted to 12 μl and 525 mM NaCl, 52.5 mM sodium citrate, 0.25% SDS. The sample was denatured at 100°C for 2 minutes and added to the array. The sample and the array were hybridized under high stringency hybridization conditions. The sample and array were covered by a 22 mm x 22 mm coverslip and placed in a humidified hybridization chamber (Corning, Acton, MA) and incubated at 65°C for 12-16 hours. Following hybridization, the arrays were washed with successive 5-minute washes in 2xSSC, 0.1%SDS; 1xSSC; and 0.1xSSC. After the final wash, the arrays were dried by centrifugation at 20xg and scanned using the GMS 418 Array Scanner (Affymetrix/Genetic Microsystems, Woburn, MA).

Example 27

Array Scanning and Data Analyses

Scanned arrays were saved as 16-bit TIFF files and analyzed using Biodiscovery's Imagene software (Los Angeles, CA). Data mining and clustering analysis was performed using Biodiscovery's GeneSight software. Prior to data analysis, all cell culture samples were normalized using all spots on the array. All mouse tissue samples were normalized to the housekeeping genes on the mouse array. Local background was calculated for each individual spot and any spot with a signal intensity less than 3 times over background or that had poor morphology was excluded from the data analyses. Only differential expression values of greater than 1.7 were considered significant. All data mining and clustering analysis performed using GeneSight was on expression ratio data that was transformed by taking the natural log (ln) of all values and normalized by Z-score. The data is transformed because of the non-Gaussian distribution of the expression ratio values. Because the ratios are bounded on the lower limit by zero, a non-Gaussian distribution is normally observed. To allow for additional statistical manipulations, the data was transformed for a more uniform distribution. The Z-score normalization method involved subtracting the mean from every observation and dividing by its standard deviation, effectively normalizing each spot to all other spots on the array.

Example 28

Sample Hybridization to Mitochondrial Array

Control cDNA samples were prepared from mRNA isolated from the LM(TK) - cell line and labeled with the Cy5 dye. Each experimental mRNA sample was labeled with the Cy3 dye, combined with the Cy5 control sample and the mixture used to hybridize the array. A representative image of a hybridized array is shown in FIG 2. Any spot on an array that appeared red was due to hybridization of a large proportion of the Cy5-labeled control LM(TK) - sample and any sample that was green was due to the hybridization of a large proportion of the Cy3-labeled experimental sample. Any spot that is yellow is an about equal co-hybridization of the two targets. The fluorescence ratio was quantitated for each spot, permitting calculation of the relative abundance of each gene's mRNA in the two samples.

Example 29

Normalization

The two fluorescent dyes that were used to label the cDNA produced during the reverse transcription of the mRNA have different structures and different emission maxima. Therefore, the two images that represent the hybridization of each of the fluorescently labeled samples were normalized to each other to account for the differences in dye behavior prior to calculating the expression ratios between the two images. One image was normalized to the other by averaging all of the spots in each image to derive a constant that was then applied to each spot. Alternatively, a predetermined set of genes that were expressed equally in the two samples under all conditions could have been used. The expression ratios of these genes were used to calculate a constant that was then applied to all spots on the array. A set of 25 housekeeping genes in Table 2 was included on a mouse array for normalization and both of these methods were used in the analysis of the mouse cell line and tissue samples. Housekeeping gene expression in the cultured cells was much more variable than in the tissue samples. Because of the variability in the housekeeping gene expression patterns in the cell line samples, normalization was done using all of the spots on the array. The expression of the housekeeping genes was much more consistent in the tissue samples and normalization using either the housekeeping genes or the average of all of the genes gave similar results.

Example 30

Clones useful for making control probes for the arrays of this invention are listed in Table 6. Sequences of the genes useful for making the control probes are provided in the sequence listings hereof.

Table 6

SEQ ID NO.	Gene Name	Complete Gene Name	GenBank Accession No.
3041	Beta Actin	Human beta actin	X63432
3042	Plant control CAB	Arabidopsis photosystem I chlorophyll a/b binding protein	X56062
3043	Plant control rbcL	Arabidopsis ribulose-1,5-bisphosphate carboxylase/oxygenase large subunit	U91966
3044	Plant control RUBISCO	Arabidopsis RUBISCO activase	X14212

It will be appreciated by those of ordinary skill in the art that samples, sample collection techniques, sample preparation techniques, probes, probe generation techniques, genes involved in mitochondrial biology, hybridization techniques, array printing techniques, physiological conditions, cell lines, mutant strains, organisms, tissues, solid substrates, and methods of data analyses other than those specifically disclosed herein are available in the art and can be employed in the practice of this invention. All art-known functional equivalents are intended to be encompassed within the scope of this invention.

REFERENCE TO SEQUENCE LISTINGS

Tables 3-5 list sequence information on the clones that are useful for making probes for practicing the methods of this invention. Clone identification numbers are usually from NIA (National Institutes of Aging, National Institutes of Health, Bethesda, MD, USA), ResGen Invitrogen (Carlsband, CA, USA) or IMAGE Consortium, LLNL (Livermore, CA, USA). Gene names and descriptions are provided for the gene interrogated by a probe made from the corresponding clone. GenBank Accession Number and Unigene Cluster ID are provided where available. The functions of certain genes are included in Table 4. Sequences of the 5' and 3' ends of the clones listed in Tables 3-4 are provided when available. If no 5' or 3' sequence was available, gene sequence from the GenBank Accession No. provided for that clone is listed in some cases. The GenBank sequence may be larger than the sequence of the clone. The instant invention may be practiced without the sequence information provided

herein using the clones or GenBank listings. Other sequences derived from the genes interrogated by probes generated from clones listed in Tables 3-5 are useful for making equivalent probes using information known in the art, i.e., unique segments of such genes may be used.

The sequence listings that correspond to the clones listed in Table 3, covering human probes SEQ ID NOS:1-994, contain information including: the sequence identification number; a GenBank Accession No. corresponding to the listed sequence or the gene interrogated by the probe containing the listed sequence; another GenBank Accession No. in parentheses which is associated with the listed sequence in Table 3; a Research Genetics (ResGen Invitrogen, Carlsbad, CA, USA) Clone ID No. identifying the clone from which the sequence was derived; the name of the gene from which the clone was derived; a description of the gene; the Unigene Cluster ID No. of the gene; the IMAGE Clone ID No., which is often the same as the ResGen Clone ID No., and information in parentheses identifying the sequence as 5' or 3' of the clone; the length of the insert of the clone; the source of the clone; the type of clone, such as cDNA; and the nucleic acid sequence.

Sequence listings for control probes are provided as SEQ ID NOS:3041-3044.

The sequence listings that correspond to the clones listed in Table 4, covering mouse probes SEQ ID NOS:995-3040, contain information including: the sequence identification number; a GenBank Accession No. corresponding to the listed sequence; the 5' and/or 3' sequence of the corresponding clone, or the gene from which the corresponding clone was derived; the name and description of the gene from which the corresponding clone was derived; the Unigene Cluster ID No. of the gene from which the corresponding clone was derived; the name of the clone from which the instant sequence was derived; additional description of the gene; a set of titles usually including Clone Name, Rearray Sequence, Parent Sequence, Other EST, and Blast Link; a list of names including, in order of the abovementioned titles, the name of the clone from which the sequence was derived, the name of the sequence with a suffix identifying it as the 5' (-5) or 3' (-3) sequence of the clone, the name of the parent sequence, and the name of another EST (expressed sequence tag), if it exists, which would be the other of the 3' or 5' sequence; the length of the sequence provided; and the nucleic acid sequence.

SEQ ID NOS:1-3044 are submitted herewith on a compact disk in file "Sequence Listing.txt" which is incorporated herein by reference.

CLAIMS

We claim:

- 1. An array comprising at least two isolated nucleotide molecules, each molecule having a sequence capable of uniquely hybridizing to a nucleic acid molecule which is an expression product of a gene involved in mitochondrial biology.
- 2. An array comprising two or more isolated nucleic acid molecules or spots, each spot comprising a plurality of isolated nucleic acid molecules, each molecule having a sequence consisting essentially of a sequence selected from the group consisting of the sequences of human probe set #1, SEQ ID NOS: 1 to 994, or mouse probe set #2, SEQ ID NOS: 995 to 3040, and sequences having at least 70% homology to the foregoing sequences.
- 3. The array of claim 2 printed on a glass slide.
- 4. The array of claim 2 comprising more than about ten spots.
- 5. The array of claim 2 comprising more than about twenty-five spots.
- 6. The array of claim 2 comprising all of the isolated nucleic acid molecules having the sequences of human probe set #1, SEQ ID NOS: 1 to 994.
- 7. The array of claim 2 comprising all of the isolated nucleic acid molecules having the sequences of mouse probe set #2, SEQ ID NOS: 995 to 3040.
- 8. The array of claim 2 also comprising one or more spots comprising control nucleic acid molecules, SEQ ID NOS:3041-3044.
- 9. A method for determining an expression profile of a sample containing nucleic acid comprising:
 - a) providing the sample;
 - b) providing an array of claim 2;
 - c) contacting said array with said sample under conditions allowing selective hybridization; and

d) measuring hybridization of nucleic acid in said sample to said array to produce an expression profile.

- 10. The method of claim 9 wherein said sample is from a mouse or a human.
- 11. A method for determining an expression profile of a first labeled sample containing nucleic acid relative to a second, differently labled sample containing nucleic acid comprising:
 - a) providing the first labeled sample;
 - b) providing the second, differently labeled sample;
 - c) providing an array of claim 2;
 - d) contacting the array with the first sample and the second sample under conditions allowing selective hybridization;
 - e) measuring hybridization of said first and said second samples to said array; and
 - f) comparing the hybridization of said first sample to the hybridization of said second sample to produce an expression profile.
- 12. The method of claim 11 wherein said second sample is a reference or a standard.
- 13. A method for determining an expression profile diagnostic of an energy-metabolism-related physiological condition comprising:
 - a) providing a labeled first sample from a first group of one or more individuals with said physiological condition;
 - b) providing a differently labeled second sample from a second group of one or more individuals without said physiological condition;
 - c) providing an array of claim 2;
 - d) contacting the array with the first sample and the said second sample under conditions allowing selective hybridization;
 - e) measuring hybridization of said first and said second samples to said array; and
 - f) comparing the hybridization of said first sample to the hybridization of said second sample to produce an expression profile diagnostic of said physiological condition.

- 14. A method of making an array comprising:
 - a) providing a prepared substrate; and
 - b) printing two or more spots in known positions on said substrate, each spot comprising a plurality of isolated nucleic acid molecules, each molecule having a sequence consisting essentially of a sequence selected from the group consisting of human probe set #1, SEQ ID NOS: 1 to 994, mouse probe set #2, SEQ ID NOS: 995 to 3040, and sequences having at least 70% homology to the foregoing sequences.
- 15. The method of claim 14 wherein said array comprises all of said isolated nucleic acid molecules in human probe set #1, SEQ ID NOS: 1 to 994.
- 16. The method of claim 14 wherein said array comprises all of said isolated nucleic acid molecules in mouse probe set #2, SEQ ID NOS: 995 to 3040.
- 17. A method of diagnosing a first individual with Complex IV Leigh's Syndrome comprising detecting in a first sample from said first individual at least about a 1.7-fold decrease in the amount of expression of genes comprising ND4, NDL4, ND6, SURF-1, SOD2, 70kD heat shock protein, VDAC4, ANT2, and glutathione peroxidase 3 compared to the amount of expression of said genes in a second sample from a second individual without Complex IV Leigh's Syndrome.
- 18. A library of at least two isolated nucleic acid molecules, each molecule having a sequence consisting essentially of a sequence selected from the group consisting of human probe set #1, SEQ ID NOS: 1 to 994, mouse probe set #2, SEQ ID NOS: 995 to 3040, and sequences having at least 70% homology to the foregoing sequences.
- 19. An array comprising at least two spots, each spot comprising a plurality of isolated nucleic acid molecules, each molecule comprising a sequence with at least 70% homology to a sequence selected from the group consisting of human probe set #1, SEQ ID NOS: 1 to 994.

20. An array comprising at least two spots, each spot comprising a plurality of isolated nucleic acid molecules, each molecule comprising a sequence with at least 70% homology to a sequence selected from the group consisting of mouse probe set #2, SEQ ID NOS: 995 to 3040.

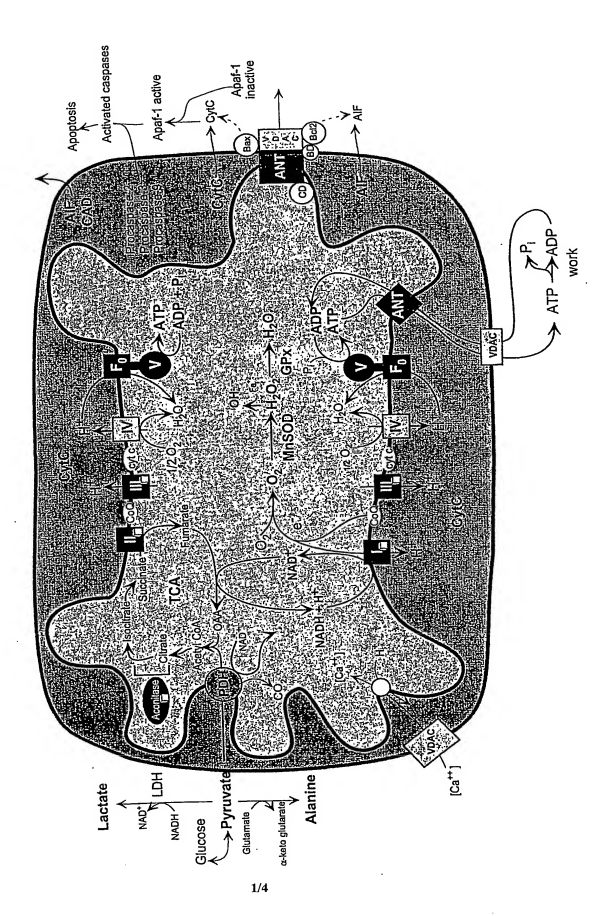


FIG. 2

